SEARCH REQUEST FORM

1.737

| SEARCH REQUEST FORM (2/3) |
|--|
| Requestor's Name: PHILLIP GAMBEL Serial Number: 08/487283 Date: 1/29/97 Phone: 308.3997 Art Unit: 1806 |
| Search Topic: Please write a detailed statement of search topic. Describe specifically as possible the subject matter to be searched. Define any terms that may be a special meaning. Give examples or relevent citations, authors, keywords, etc., if known. For sequences, please anoth a copy of the sequence. You may include a copy of the broadest and/or most relevent claim(s). |
| sep/interference :search |
| SEQ 10 NO. 1 + 2 - |
| REQUEST JUN |
| There |
| |
| |
| |
| 1/28/98 STAFF USE ONLY |
| Date completed: //26/76 Search Site Vendors Searcher: STIC IG MC |
| Terminal time: CM-1 STN Elapsed time: Pre-S Dialog |
| Elapsed time: Pre-S Dialog CPU time: APS |
| Total time: N.A. Sequence Geninfo |
| Number of Scarches: A.A. Sequence SDC |
| Number of Databases: Structure DARC/Questel |

_ Bibliographic

_ Other

Jan 28 12:17

US-08-487-283A-1.mm

(TM)

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 University of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

protein - protein database search, using Smith-Waterman algorithm MPsrch pp

Wed Jan 28 12:10:17 1998; MasPar time 3.57 Seconds 179.105 Million cell updates/sec Run on:

Tabular output not generated.

Title:

>US-08-487-283A-1 (1-21) from US08487283A.pep 141 1 VIDHQGTKSSKCVRQKVEGSS 21 Description: Perfect Score:

Sequence:

PAM 150 Gap 15 Scoring table:

95051 seqs, 30469580 residues

Searched:

Post-processing: Minimum Match 0% Listing first 45 summaries

Database:

1:ann1 2:ann2 3:ann3 4:ann4 5:unann1 6:unann2 7:unann3 8:unann4 9:unann5 10:unann6 11:unann7 12:unann8 13:unann9 14:unann10 15:unann11 16:unann12 17:unenc

Mean 25.513; Variance 35.018; scale 0.729 Statistics: Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| Query Score Match Length DB | uery atch L | - A (| angth | 88 | 10 | Description | Pred. No. |
|--------------------------------|----------------|---------|-------|----|---------|-----------------------|------------|
| | | 1676 | | ٠. | С5НО | complement C5 precur | 3.46e - 20 |
| | | 1680 2 | 7 | | CSMS | complement C5 precur | 9.17e-03 |
| | | 213 11 | 7 | | S76070 | hypothetical protein | • |
| 59 41.8 940 10 | | 940 10 | 2 | _ | S49087 | lactoferrin binding | 9.30e-01 |
| | | 1002 15 | 15 | | A56678 | yemanuclein-alpha - | • |
| | | 205 16 | 16 | | 149364 | protein tyrosine pho | |
| | | 223 16 | 16 | | 149365 | protein tyrosine pho | |
| 40.4 | | 117 8 | œ | | 168524 | ribosomal protein L3 | • |
| 40.4 | | 264 18 | 18 | | 828969 | N-carbamoy Isarcosine | • • |
| | | 537 13 | 13 | | B334F5 | spore coat protein S | |
| | | 540 13 | 13 | | \$21825 | vicilin-like storage | 3.42e+00 |
| | | | | | | | |

US-08-487-283A-1.rpr Jan 28 12:17

C)

| qlobulin-15, GLB1S - 3.42e+00 | transposable element 5.22e+00 | spore coat peptide C 7.93e+00 | | P element - fruit fl 7.93e+00 | dipoptidyl aminopept 7.93e+00 | -w | E4 11K protein 1. | 11K | e-rich | epithelin/granulin p 1.20e+01 | omega-crystallin - g 1.20e+01 | CO | platelet-derived gro 1.20e+01 | TMV resistance prote 1.20e+01 | cell-fate determinin 1.20e+01 | Glb1-0 protein - mai 1.80e+01 | globulin-10, GLB10 - 1.80e+01 | VSG expression site- 1.80e+01 | ornithine cyclodeami 1.80e+01 | methylaspartate ammo 1.80e+01 | vicilin-like storage 1.80e+01 | 1. | hypothetical protein 2.69e+01 | protein 2. | drogenas 2. | pyruvate kinase (EC 2.69e+01 | major envelope glyco 2.69e+01 | | ein 2. | protein-tyrosine kin 2.69e+01 | protein-tyrosine kin 2.69e+01 | retrovirus-related p 2.69e+01 | clathrin heavy chain 2.69e+01 |
|-------------------------------|-------------------------------|-------------------------------|--------|-------------------------------|-------------------------------|--------|-------------------|--------|--------|-------------------------------|-------------------------------|--------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|--------|-------------------------------|------------|-------------|------------------------------|-------------------------------|--------|--------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| A53234 | A32494 | E47119 | G64383 | 546281 | A30107 | S57380 | Q4ADE5 | Q4ADE2 | 541772 | C38128 | 543428 | 154205 | PFHUGA | A54810 | A49128 | 521823 | C53234 | VMUT21 | DUAGO | S55582 | B53234 | S00954 | 523314 | 507237 | A35260 | S29004 | VGNVAC | A37197 | 523319 | A47511 | JC4127 | • | S52588 |
| 13 | 12 | 12 | 12 | 18 | - | 14 | 4 | 4 | _ | œ | 14 | 15 | | 8 | 16 | 13 | 13 | ٣ | 7 | 9 | 13 | 12 | 6 | 6 | 2 | S | 4 | œ | 6 | 16 | 16 | 9 | œ |
| 573 | 775 | 148 | 224 | 562 | 818 | 1030 | 116 | 116 | 148 | 589 | 591 | 699 | 1089 | 1144 | 2471 | 122 | 122 | 329 | 354 | 354 | 582 | 982 | 238 | 250 | 337 | 462 | 529 | 574 | 1084 | 1129 | 1132 | 1291 | 1678 |
| 39.7 | 39.0 | 38.3 | 38.3 | 38.3 | 38.3 | 38.3 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 36.9 | 36.9 | 36.9 | 36.9 | 36.9 | 36.9 | 36.9 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 |
| 56 | 22 | 24 | 24 | 24 | 24 | 54 | 23 | 23 | 23 | 23 | 23 | 23 | 23 | 23 | 23 | 25 | 25 | 25 | 25 | 25 | 25 | 25 | 51 | 21 | 51 | 21 | 51 | 51 | 21 | 21 | 21 | 21 | 21 |
| 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 52 | 56 | 27 | 28 | 29 | 30 | 31 | 32 | 33 | 34 | 32 | 36 | 37 | 38 | 39 | 40 | 41 | 42 | 43 | 44 | 45 |

ALIGNMENTS

| C5HU #tvpe complete | complement C5 precursor - human | C5a anaphylatoxin; C5b | formal_name Homo sapiens fcommon_name man | 30-Sep-1992 #sequence_revision 30-Sep-1992 #text_change 27-Feb-1997 | A40075; A27689; A01267; A01266; S15121 | A40075 | Haviland, D.L.; Haviland, J.C.; Fleischer, D.T.; Hunt, A.; | Wetsel, R.A. | J. Immunol. (1991) 146:362-368 | Complete cDNA sequence of human complement pro-C5. Evidence | of truncated transcripts derived from a single copy gene. | <pre>#cross-references MUID:91079575</pre> | A40075 | ##molecule_type_mRNA | ues 1-1676 ##label HAV | ##cross-references GB:M57729 | 518-Ser was also found | A27689 | Wetsel, R.A.; Lemons, R.S.; Le Beau, M.M.; Barnum, S.R.; | Noack, D.; Tack, B.F. | Biochemistry (1988) 27:1474-1482 | Molecular analysis of human complement component C5: | localization of the structural gene to chromosome 9. |
|---------------------|---------------------------------|------------------------|---|--|--|-----------|--|--------------|--------------------------------|---|---|--|------------|----------------------|------------------------|------------------------------|------------------------|-----------|--|-----------------------|----------------------------------|--|--|
| RESULT 1 ENTRY | TITLE | CONTAINS | ORGANISM | DATE | ACCESSIONS | REFERENCE | #authors | | # journal | #title | | #cross-ref | faccession | ##molec | ##residues | ffcross | ##note | REFERENCE | #authors | | # journal | #title | |

```
A35530; A27538; A40429
A35530
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     C5a anaphylatoxin; C5b
                                                                                                                                                                                                                                                                                                                                                                                                                                 experimental/
                                                                                                                                                                                                                                                                                                                                                                                  experimental/
  #label C5A\
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              872 vidhqqtksskcvrqkvegss 892
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        1 VIDHQGTKSSKCVRQKVEGSS 21
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            cross-references MUID:87185363
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       #cross-references GB:J05234
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  predicted
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity 100.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                21; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             16-Feb-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         complement
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A35530
                                                                                                                                                     111–732, 866–1527,
                                                                                                      567-810, 634-669,
                                                                                                                             598-724, 699-731
                                                                                                                                                                                                                                                                                                                                                                                                                                                             911, 1115, 1630
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ##residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     #accession
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            #accession
                                                                                                                                                                                                   1375-1505,
                                                                                                                                                                                                                               1405-1474,
                                                                                                                                                                                                                                                                              1532-1606,
                                                                                                                                                                                                                                                                                                        1553-1676,
                                                                                                                                                                                 1101 - 1159
                                                                                                                                                                                                                                                       1520-1525
                                                                                                                                                                                                                                                                                                                                 1654-1657
                                                                          752-1676
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 #journal
#title
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               #authors
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       authors
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        f journal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          #authors
                                                                                                                                                                                                                                                                                                                                                                                                          751-752
                         678-751
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Matches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ACCESSIONS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                #title
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CONTAINS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ORGANISM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SUMMARY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 쇰
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ò
                                                                                                                                                Fernandez, H.N.; Hugli, T.E.
J. Biol. Chem. (1978) 253:6955-6964
Primary structural analysis of the polypeptide portion of
human C5a anaphylatoxin. Polypeptide sequence determination
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     complement alternate pathway; complement pathway; cytolysis; glycoprotein; inflammation; membrane attack complex; plasma
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Isolation and sequence analysis of a cDNA clone encoding the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            removal of four basic residues. C5 convertase releases C5a anaphylatoxin from the amino end of the alpha chain, generating
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Activation of C5 initiates the spontaneous assembly of the late complement components, C5-C9, into the membrane attack complex. C5b has a transient binding site fcr C6. The C5b-C6 complex is the foundation upon which the membrane attack complex is
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Bohnsack, J.F.; Mollison, K.W.; Buko, A.M.; Ashworth, J.C.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Group B streptococci inactivate complement component C5a by
                                                                                                                                                                                                                                                       and assignment of the oligosaccharide attachment site in
                                                                                                                                                                                                                                                                                                                                                                                                                               Lundwall, A.B.; Wetsel, R.A.; Kristensen, T.; Whitehead,
A.S.; Woods, D.E.; Ogden, R.C.; Colten, H.R.; Tack, B.F.
J. Biol. Chem. (1985) 260:2108-2112
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              #domain signal sequence #status predicted #label SIG\
#product complement C5 #status predicted #label MAT\
#product C5b #status predicted #label C5B\
#product complement C5 and C5b beta chain #status
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         product complement C5 alpha chain *status predicted
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              'SLALSPRLECNGKISGHCKLRLPGSSDSPASASQVAGITGTHHHAQPT'
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       the carboxyl-terminal part of the sequence in this report appears to be derived from translation of
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Complement C5 contains two disulfide-linked chains, formed by
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     C5a has potent spasmogenic and chemotactic activity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               enzymic cleavage at the C-terminus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              #superfamily alpha-2-macroglobulin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Biochem. J. (1991) 273:635-640
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     fifth complement component.
#cross-references MUID:85130937
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                predicted | | label C5BB\
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ALU repeat sequence
                                                ##molecule_type mRNA
##residues 412-1676 ##label WET
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  C5b (beta and alpha' chains).
                                                                                                                                                                                                                                                                                                                                               ##molecule_type protein
##roaidues 678-751 ##label FER
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ##label LUN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ##cross-references GDB:119734
                                                                                                    ##cross-references GB:M18879
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    #cross-references MUID:91144547
tross-references MUID:88209511
                                                                                                                                                                                                                                                                                                        fcross-references MUID:79005687
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ##cross-references GB:K02874
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     412-854,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  annotation
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           #molecule_type mRNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  #map_position 9q33-9q33
CLASSIFICATION #superfam
                                                                                                                                                                                                                                                                                                                                                                                                       A01266
                                                                                                                                A01267
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A01266
                         A27689
                                                                                                                                                                                                                                                                                                                                 A01267
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        GDB:C5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           assembled
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     19-673, 678-1676
19-673, 752-1676
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            #residues
                           #accession
                                                                                                                                                                                                                                                                                                                                      #accession
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           #accession
                                                                                                                                                                                                                                                                                                                                                                                                                                      #authors
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  #contents
                                                                                                                                                       authors
                                                                                                                                                                                 # journal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      # journal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           #authors
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              # journal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         678-1676
                                                                                                                                REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                              REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    REFERENCE
```

#dene GENETICS

COMMENT

COMMENT

KEYWORDS

Jan 28 12:17

#product C5b alpha' chain #status predicted #label C5BA\ Haviland, D.L.; Haviland, J.C.; Fleischer, D.T.; Wetsel, R.A. J. Biol. Chem. (1991) 266:11818-11825 Structure of the murine fifth complement component (C5) gene. ö A large, highly interrupted gene with a variant donor splice site and organizational homology with the third and #journal J. Biol. Chem. (1990) 265:2435-2440

#title Deficiency of the murine fifth complement component (C5). A 2-base pair gene deletion in a 5'-exon.

#cross-references MUID:90153853 !product C5a anaphylatoxin #status experimental #label #binding site carbohydrate (Asn) (covalent) #status #binding_site carbohydrate (Asn) (covalent) #status Gaps #formal name Mus musculus #common name house mouse 19-Nov-1988 #sequence_revision 15-Oct-1994 #text_change #length 1676 #molecular-weight 188330 #checksum 3858 #cleavage_site Arg-Leu (C5 convertase) #status Primary structure of the fifth component of murine ö 100.0%; Score 141; DB 2; Length 1676; 100.0%; Pred. No. 3.46e-20; Wetsel, R.A.; Fleischer, D.T.; Haviland, D.L. 0; Indels disulfide bonds fstatus predicted/ Wetsel, R.A.; Ogata, R.T.; Tack, B.F. 0; Mismatches complement C5 precursor - mouse Biochemistry (1987) 26:737-743 #molecule_type mRNA #residues 1-215,'L' ##label WET #type complete

US-98-487-283A-1.pp

ö Nakamura, Y.; Miyajima, N.; Hirosawa, M.; Sugiura, M.; Sasamoto, S.; Kimura, T.; Hosouchi, T.; Matsuno, A.; Muraki, A.; Nakazaki, N.; Naruo, K.; Okumura, S.; Shimpo, the nucleotide sequence was submitted to the EMBL Data Poolman, X.Y.Z.; Tommassen, X.Y.Z. submitted to the EMBL Data Library, June 1994 Molecular charactarization of the structural gene for the iacto-ferrin receptor of the meningococcal strain H44/76 Kaneko, T.; Sato, S.; Kotani, H.; Tanaka, A.; Asamizu, E.; Pettersson, A.M.; Klarenbeek, X.Y.Z.; van Deurzen, X.Y.Z.; Sequence analysis of the genome of the unicellular cyanobacterium Synechocystis sp. PCC6803. II. Sequence determination of the entire genome and assignment of S.; Takeuchi, C.; Wada, T.; Watanabe, A.; Yamada, M.; Yasuda, M.; Tabata, S. DNA Res. (1996) 3:109-136 Gaps 25-Apr-1997 #sequence_revision 25-Apr-1997 #text_change #formal name Neisseria meningitidis
16-Feb-1995 #sequence_revision 12-May-1995 #text_change #length 213 #molecular-weight 22745 #checksum 2191 lactoferrin binding protein - Neisseria meningitidis ö Length 213; 6; Indels Score 62; DB 11; 1 Pred. No. 2.43e-01; potential protein-coding regions 7; Mismatches #type complete Library, June 1996 1-213 ##label KAN ##cross-references EMBL:D63999 92 ilayigakasdcikpkvvss 111 preliminary preliminary 44.0%; Best Local Similarity 35.0%; 7; Conservative 25-Apr-1997 12-May-1995 ##molecule_type DNA 574322 S76070 576070 S49087 549087 S49087

ö

Gaps

ö

Indels

5; Mismatches

Length 940;

Score 59; DB 10; 1 Pred. No. 9.30e-01;

1-940 ##label PET

US-08-487-283A-1.mm

Jan 28 12:17

A single mutation converts a novel-phosphotyrosine binding ribosomal protein L34 - human #formal name Homo sapiens #common_name man 24-May-1996 #sequence_revision 24-May-1996 #text_change #formal name Arthrobacter sp.
19-Mar-1997 #sequence_revision 19-Mar-1997 #text_change
19-Mar-1997 #formal name Mus musculus #common name house mouse
02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change Wishart, M.J.; Denu, J.M.; Williams, J.A.; Dixon, J.E. J. Biol. Chem. (1995) 270:26782-26785 Rommens, J.M.; Durocher, F.; McArthur, J.; Tonin, P.; Generation of a transcription map at the HSD17B locus ##molecule_type mRNA
#fresidues 1-117 ##label RES
#fresidues 1-117 ##label RES
#fross-references GB:L38941; NID:q1008855; CDS_PID:q1008856
CLASSIFICATION #prefamily rat ribosomal protein L34
SUMMARY #length 117 #molecular-weight 13305 #checksum 4392 ffcross-references EMBL:U34973; NID:g1063624; CDS_PID:g1063626 N-carbamoylsarcosine amidohydrolase (EC 3.5.1.59) Arthrobacter sp. #length 223 #molecular-weight 25416 #checksum preliminary; translated from GB/EMBL/DDBJ Query Match 41.1%; Score 58; DB 16; Length 223; Best Local Similarity 25.0%; Pred. No. 1.44e+00; preliminary; translated from GB/EMBL/DDBJ domain into a dual-specificity phosphatase 40.4%; Score 57; DB 8; Length 117; Pred. No. 2.22e+00; Mismatches 9; Mismatches centromeric to BRCA1 at 17q2] Genomics (1995) 28:530-542 #type complete #type complete 1-223 ##label 51 ilqkhgithiicirqniean 70 1 VIDHQGTKSSKCVRQKVEGS 20 Best Local Similarity 41.7%; 92-Jul-1996 16-Feb-1997 5; Conservative 5; Conservative Leblanc, J. ##molecule_type mRNA 6 GTKSSKCVRQKV 17 80 gsmcakcvrdri 91 168524 154209 149364 149365 168524 I 68524 149365 528969 ##residues #status ##status *accession Query Match #accession Query Match #introns authors journal #authors | journal **ACCESSIONS** ACCESS IONS ftitle REFERENCE Matches #title Matches REFERENCE ORGANISM GENETICS ORGANISM ORGANISM SUMMARY RESULT ENTRY RESULT ENTRY TITLE TITLE DATE DATE 쇰 ð 쇰 8 ö ö The yemanuclein-alpha: a new Drosophila DNA binding protein specific for the occyte nucleus. A56678 A single mutation converts a novel-phosphotyrosine binding #formal name Drosophila melanogaster 08-Jul-1995 #text_change 16-Peb-1997 Gaps Gaps protein tyrosine phosphatase - mouse #formal name Mus musculus #common name house mouse 02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change A56678 #type complete yemanuclein-alpha - fruit fly (Drosophila melanogaster) DNA binding; oocyte flength 1002 fmolecular-weight 109310 fchecksum 4278 Wishart, M.J.; Denu, J.M.; Williams, J.A.; Dixon, J.E. J. Biol. Chem. (1995) 270:26782-26785 ##cross-references EMBL:U34973; NID:q1063624; CDS PID:q1063625 Y #length 205 #molecular-weight 23683 #checksum 2745 ö ö Ait-Ahmed, O.; Bellon, B.; Capri, M.; Joblet, C.; Query Match 41.8%; Score 59; DB 15; Length 1002; Best Local Similarity 50.0%; Pred. No. 9.30e-01; Query Match 41.1%; Score 58; DB 16; Length 205; Best Local Similarity 25.0%; Pred. No. 1.44e+00; 5; Mismatches 1; Indels domain into a dual-specificity phosphatase 149364 6; Indels Mismatches ##cross-references FlyBase:FBgn0005596 trons 80/3; 154/3; 428/1 477/2; 557/2 Thomas-Delaage, M. Mech. Dev. (1992) 37:69-80 #type complete 1-1002 ##label AIE ##cross-references GB:X63503 preliminary 51 ilqkhgithiicirqniean 70 1 VIDHQGTKSSKCVRQKVEGS 20 FlyBase:yem&agr A56678; S22146 02-Jul-1996 6; Conservative 5; Conservative ##molecule_type DNA 47 tktakcirikld 58 A56678 149364 149364

149364

Pauthors

ACCESSIONS

REFERENCE

TITLE ORGANISM

DATE

RESULT

ENTRY

| journal

#title

#accession

Query Match

SUMMARY

Matches

쇰 8

ö

Gaps

ö

1; Indels

149365

RESULT

ENTRY

ö

Gaps

ö

6; Indels

##residues

#introns

KEYWORDS

SUMMARY

gene

GENETICS

Matches

g ð

accession

#authors

ACCESSIONS

ORGANISM

ENTRY

REFERENCE

#journal

title

US-08-487-283A-1.rpr

Jan 28 12.17

9

ö ö Gaps globulin-15, GLBIS - maize
#formal name Zea mays #common_name maize
02-May-1994 #sequence_revision 18-Nov-1994 #text_change
05-Apr-1995 Gaps Belanger, F.C.; Kriz, A.L. Genetics (1991) 129:863-872 Molecular basis for allelic polymorphism of the maize 170/1; 195/2; 222/2; 319/2 #length 540 #molecular-weight 60239 #checksum 1419 sequence extracted from NCBI backbone #length 573 #molecular-weight 65075 #checksum 3569 ö ö Score 56; DB 13; Length 540; Pred. No. 3.42e+00; 2; Indels Score 56; DB 13; 1 Pred. No. 3.42e+00; Mismatches Mismatches #fexperimental source inbred line Va 26 #type complete 1-540 ##label KRI ##cross-references EMBL:X59084 Globulin-1 gene. \$\frac{4}{cross-references}\$ MUID:92090707 preliminary preliminary Query Match 39.7%; Best Local Similarity 58.3%; Query Match 39.7%; Best Local Similarity 58.3%; 7; Conservative 7; Conservative 32 hgghksgrcvrr 43 ##molecule_type DNA 4 HQGTKSSKCVRQ 15 32 hgghksgrcvrr 43 4 HQGTKSSKCVRQ 15 ##molecule_type DNA Glb1-S A53234 A53234 ##residues ##status ##status #accession #accession ##note #introns 12 authors journal **ACCESSIONS** Matches #title Matches # dene REFERENCE GENETICS ORGANISM SUMMARY

#formal name Xenopus laevis #common name African clawed frog 12-Oct-1989 #sequence_revision 31-Dec-1993 #text_change

31-Dec-1993

preliminary

fstatus

#submission

transposable element Txlc protein 1 - African clawed frog

#type complete

A32494

13

1-775 ##label GAR

#molecule_type DNA

##residues

##cross-references GB:M26915

#note

SUMMARY

US-08-487-283A-1.mr

K.M.; Hurst, M.A.; Kaine, B.P.; Borodovsky, M.; Klenk, H.P.; Fraser, C.M.; Smith, H.O.; Woese, C.R.; Venter, J.C. Science (1996) 273:1058-1073

Complete genome sequence of the methanogenic archaeon, Methanococcus jannaschii.

the authors translated the codon ATT for residue as Gln, and AAG for residue 288 as Leu #length 775 #molecular-weight 82355 #checksum 6734

preliminary; nucleic acid sequence not shown; G64383 #accession

translation not shown ##status

ö

Gaps

ö

3; Indels

5; Mismatches

7; Conservative

Matches

617 sntskcvssevegtp 631

염 8

7 TKSSKCVRQKVEGSS 21

Query Match 39.0%; Score 55; DB 15; Length 775; Best Local Similarity 46.7%; Pred. No. 5.22e+00;

#map position REV597638-596964 GENETICS

Query Match 38.3%; Best Local Similarity 33.3%;

spore coat peptide Cotz - Bacillus subtilis #formal name Bacillus subtilis 21-Sep-1993 #sequence_revision 18-Nov-1994 #text_change 18-Nov-1994

#start_codon
KEYWORDS

hydrolase #length 224 #molecular-weight 25037 #checksum 2215 SUMMARY

ö Score 54; DB 12; Length 224; Pred. No. 7.93e+00; 4; Mismatches 8; Indels 6; Conservative Matches

118 iledmgvevvkcgrgkvd 135

1 VIDHQGTKSSKCVRQKVE 18 ð Search completed: Wed Jan 28 12:10:33 1998 Job time : 16 secs.

##molecule_type DNA

Gaps

ö

polypeptides present in the insoluble fraction of the spore coat of Bacillus subtilis.

#cross-references MUID:93285989
#facession E47119

Zhang, J.; Fitz-James, P.C.; Aronson, A.I. J. Bacteriol. (1993) 175:3757-3766

A47119

authors journal

ACCESSIONS

ORGANISM

DATE

RESULT

ENTRY TITLE REFERENCE

ö

Gaps

..

2; Indels

Mismatches

7; Conservative

Matches

염 ð

4 ktsscvreave 14

15

RESULT ENTRY

TITLE

#length 148 #molecular-weight 16534 #checksum 4681

sequence extracted from NCBI backbone

##cross-references NCBIN:133538; NCBIP:133548

##note

SUMMARY

##status preliminary ##molecule_type_nucleic_acid ##residues 1-148 ##label_ZHA

Score 54; DB 12; Length 148; Pred. No. 7.93e+00;

Query Match 38.3%; Best Local Similarity 63.6%;

Bult, C.J.; White, O.; Olsen, G.J.; Zhou, L.; Fleischmann, R.D.; Sutton, G.G.; Blake, J.A.; FitzGerald, L.M.; Clayton, R.A.; Gocayne, J.D.; Kerlavage, A.R.; Dougherty, B.A.; Tomb, J.F.; Adams, M.D.; Reich, C.I.; Overbeek, R.; Kirkness, E.F.; Weinstock, K.G.; Merrick, J.M.; Glodek, A.; Scott, J.L.; Geoghagen, N.S.M.; Weidman, J.F.; Fuhrmann, J.L.; Nguyen, D.; Utterback, T.R.; Kelley, J.M.; Peterson, J.D.; Sadow, P.W.; Hanna, M.C.; Cotton, M.D.; Roberts,

G64383 #type complete riboflavin-specific deaminase (EC 3.5.4.-) - Methanococcus

13-Sep-1996 #sequence_revision 13-Sep-1996 #text_change

13-Sep-1996

A64300

#authors

ACCESSIONS

REFERENCE

formal name Methanococcus jannaschii

jannaschii

ORGANISM

DATE

Jan 28 12:17

-)

US-08-487-283A-1.rsp

| ', ' |
|-----------|
| |
| |
| |
| |
| |
| |

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 University of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

Œ

protein - protein database search, using Smith-Waterman algorithm MPsrch_pp

Wed Jan 28 12:10:52 1998; MasPar time 2.26 Seconds 197.189 Million cell updates/sec Run on:

Tabular output not generated.

>US-08-487-283A-1 (1-21) from USO8487283A.pep 141 1 VIDHQGTKSSKCVRQKVEGSS 21 Description: Perfect Score: Sequence:

PAM 150 Gap 15 Scoring table:

59021 seqs, 21210388 residues Searched:

Post-processing: Minimum Match 0%

Listing first 45 summaries

swiss-prot34 Database:

1:part1 2:part2 3:part3 4:part4 5:part5 6:part6 7:part7 8:part8 9:part9 10:part10 11:part11

Mean 26.425; Variance 29.915; scale 0.883 Statistics:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| Result No. | Score | Query Match | Query Match Length DB | DB | ID | Description | Pred. No. |
|---------------|-------|----------------|--------------------------|----|------------|-----------------------|-----------|
| 1 | 141 | 100.0 | 1676 | 2 | CO5 HUMAN | COMPLEMENT C5 PRECURS | 1.16e-24 |
| 7 | 69 | 48.9 | 1680 | 7 | CO5 MOUSE | COMPLEMENT C5 PRECURS | 7.22e-04 |
| m | 59 | 41.8 | 943 | 2 | IROA NEIME | IRON-REGULATED OUTER | 1.65e-01 |
| ♥ | 59 | 41.8 | 1002 | Π | YEMA DROME | YEMANUCLEIN-ALPHA. | 1.65e-01 |
| S | 57 | 40.4 | 116 | œ | RL34 HUMAN | 60S RIBOSOMAL PROTEIN | 4.57e-01 |
| 9 | 57 | 40.4 | 264 | 7 | CSH ARTSP | N-CARBAMOYLSARCOSINE | 4.57e-01 |
| 7 | 26 | 39.7 | 537 | 6 | SP70 DICDI | SPORE COAT PROTEIN SP | 7.54e-01 |
| œ | 26 | 39.7 | 573 | 4 | GLB1 MAIZE | GLOBULIN-1 S ALLELE P | 7.54e-01 |
| 6 | 26 | 39.7 | 1357 | 00 | RPOB PSEPU | DNA-DIRECTED RNA POLY | 7.54e-01 |
| 10 | 55 | 39.0 | 775 | 1 | YTX1 XENLA | TRANSPOSON TX1 HYPOTH | 1.24e+00 |
| Ξ | 54 | 38.3 | 148 | 7 | COTZ BACSU | SPORE COAT PROTEIN 2. | 2.01e+00 |
| 12 | 54 | 38,3 | 818 | m | DAP2 YEAST | DIPEPTIDYL AMINOPEPTI | 2.01e+00 |
| 13 | 24 | 38.3 | 1039 | 1 | YR71_CAEEL | HYPOTHETICAL 118.2 KD | 2.01e+00 |

US-08-487-283A-1.rsp Jan 28 12:17

c v

| 3.25e+00 | 3.25e+00 | 3.25e+00 | 3.25e+00 | 3.25e+00 | 5.22e+00 | 5.22e+00 | 5.22e+00 | 8.32e+00 | 1.32e+01 | 1.32e+01 | 1.32e+01 | 1.32e+01 | 1.32e+01 | 1.32e+01 | 1.32e + 01 | 1.32e+01 | 1.32e+01 | 2.06e+01 | 2.06e+01 | 2.06e+01 | 2.06e+01 | 2.06e+01 |
|----------------------|----------------------|-----------------------|----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|-----------------------|----------------------|----------------------|-----------------------|-----------------------|-----------------------|------------------------|-----------------------|-----------------------|-----------------------|-----------------------|----------------------|-----------------------|-----------------------|
| PROBABLE EARLY E4 11 | PROBABLE EARLY E4 11 | GRANULINS PRECURSOR (| GALACTOCEREBROSIDASE | ALPHA PLATELET-DERIVE | VSG EXPRESSION SITE-A | ORNITHINE CYCLODEAMIN | P2X PURINOCEPTOR 5 (A | PROBABLE 60S RIBOSOMA | MAJOR ENVELOPE GLYCOP | ALCOHOL DEHYDROGENASE | PYRUVATE KINASE I (EC | MAJOR ENVELOPE GLYCOP | ESTROGEN RECEPTOR (ER | GRANULINS PRECURSOR (| TRACHEOBRONCHIAL MUCI | VITELLOGENIN 6 PRECUR | CLATHRIN HEAVY CHAIN. | 50S RIBOSOMAL PROTEIN | HYPOTHETICAL 19.7 KD | HYPOTHETICAL 22.7 KD | D-RIBOSE-BINDING PERI | D-RIBOSE-BINDING PERI | TROPONIN T, CARDIAC M | BENZENE 1, 2-DIOXYGENA | BETA-FRUCTOFURANOSIDA | PROTEIN DWNT-5 PRECUR | VARIANT SURFACE GLYCO | CHLOROPLAST 50S RIBOS | HYPOTHETICAL 30.6 KD | MAJOR ENVELOPE GLYCOP | GC-RICH SEQUENCE DNA- |
| E411 ADE05 | E411_ADE02 | GRN MOUSE | GALC HUMAN | PGDS HUMAN | ESG2 TRYBB | OCD AGRT5 | P2X5 RAT | YIF2 YEAST | VP67 NPVGM | ADH1 ZYMMO | KPY1 ECOLI | VP67 NPVAC | ESTR SALIR | GRN RAT | MUC5 HUMAN | VIT6 CAEEL | CLH DROME | RL14 HAEIN | YJB6 YEAST | Y045 NPVAC | RBSB ECOLI | RBSB SALTY | TRT2 CHICK | BEDA PSEPU | INVA DAUCA | WNT5 DROME | VSMO_TRYBB | RK14 ODOSI | YD6A SCHPO | VP67 NPVCF | GCF HUMAN |
| ٣ | m | 4 | 4 | - | m | _ | _ | Ξ | 10 | - | ა | 10 | 3 | 4 | 9 | 2 | 7 | 00 | 11 | 10 | œ | œ | 6 | | Ŋ | 10 | 10 | œ | 11 | 10 | 4 |
| 116 | 116 | 589 | 699 | 1089 | 329 | 354 | 455 | 121 | 337 | 337 | 462 | 512 | 574 | 588 | 1056 | 1651 | 1678 | 123 | 171 | 192 | 296 | 296 | 301 | 410 | 592 | 1010 | 72 | 121 | 267 | 509 | 784 |
| 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 36.9 | 36.9 | 36.9 | 36.2 | 36.2 | 36.2 | 36.5 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 36.2 | 35.5 | 35.5 | 35.5 | 35.5 | 35.5 | 35.5 | 35.5 | 35.5 | 35.5 | 34.8 | 34.8 | 34.8 | 34.8 | 34.8 |
| 53 | 23 | 23 | 23 | 23 | 25 | 25 | 25 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 51 | 51 | 51 | 20 | 20 | 20 | 20 | 20 | 20 | 20 | 20 | 20 | 49 | 49 | 49 | 49 | 49 |
| 14 | 15 | 16 | 11 | 18 | 19 | 70 | 21 | 22 | 23 | 54 | 52 | 56 | 27 | 87 | 53 | 30 | 31 | 32 | 33 | 34 | 32 | 36 | 37 | 88 | 39 | 40 | 41 | 45 | 43 | 44 | 45 |

ALIGNMENTS

ö Gaps ö Score 141; DB 2; Length 1676; 0; Indels US-08-487-283A-1.rsp 9D5c6E59 CRC32; Pred. No. 1.16e-24; 0; Mismatches POTENTIAL. POTENTIAL. POTENTIAL. · S M., 872 vidhqgtksskcvrqkvegss 892 1 VIDHQGTKSSKCVRQKVEGSS 21 1676 AA; 188331 100.0%; Best Local Similarity 100.0%; 21; Conservative 911 1115 1630 518 DISULFID Jan 28 12.17 CARBOHYD CARBOHYD CARBOHYD SEQUENCE Query Match CARBOHYD VARIANT Matches 셤 ð

RESULT

BIOCHEMISTRY 26:737-743(1987).

-!- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE SPONTANDOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9, INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIENT BINDING SITE FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYTIC COMPLEX IS ASSEMBLED.
-!- SUBMINIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE BOND: C5 CONVERTAGE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN, RELEASING C5A ANAPHYLATOXIN & GENERATING C5B (BETA CHAIN + ALPHA') EUKARYOTA; METAZOA; CHORDATA; VEKTEBRATA; TETRAPODA; MAMMALIA; EUTHERIA; RODENTIA. 01-NOV-1990 (REL. 16, LAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN). PRT; 1680 AA. WETSEL R.A., FLEISCHER D.T., HAVILAND D.L.; WETSEL R.A., OGATA R.T., TACK B.F.; J. BIOL. CHEM. 265:2435-2440(1990) 01-JAN-1988 (REL. 06, CREATED) SEQUENCE OF 41-1680 FROM N.A. STANDARD; MUS MUSCULUS (MOUSE) [1] SEQUENCE FROM N.A. 90153853 MEDLINE; 87185363 LT 2 CO5_MOUSE MEDLINE; P06684;

FUNCTION: DERIVED FROM PROTEDLYTIC DEGRADATION OF COMPLEMENT C5, C5 ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS: IT PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF POLYMORPHONUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS) SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN. -1

-!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN PROSITE; PS00477; ALPHA_2 MACROGLOBULIN. EMBL; M35525; G309124; -. EMBL; M35526; G309123; -. PIR; A27538; A27538. PIR; A35530; A35530. HSSP; P01032; 1C5A.

PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;

SIGNAL.

SIGNAL

CHAIN

COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN;

US-08-487-283A-1.rsp

Ø

Jan 28 12:17

අ ð ö

Gaps

ö

1; Indels

5; Mismatches

6; Conservative

116 AA.

PRT;

STANDARD;

Score 59; DB 11; Length 1002; Pred. No. 1.65e-01;

41.8%;

NUCLEAR LOCALIZATION SIGNAL (POTENTIAL) -! - FUNCTION: MAY PLAY A KEY ROLE IN EGG ORGANIZATION. IT MAY BE A AIT-AHMED O., BELLON B., CAPRI M., JOBLET C., THOMAS-DELAAGE M.; MECH. DEV. 37:69-80 (1992). -!- TISSUE SPECIFICITY: OCCYPE-SPECIFIC.
-!- DEVELOPMENTAL STAGE: EXPRESSED AT ALL OCGENIC STAGES.
-!- SUBCELLULAR LOCATION: NUCLEAR.
PMBL; X63503; G8838; -.
PMF; S2146; S22146.
FLYBASE; FEGN0005596; YEM-ALPHA. 698 698 S -> L (IN CDNA SEQUENCE) 1002 AA; 109310 MW; 955FDZC1 CRC32; ASP/GLU-RICH (ACIDIC). 2 X 12 AA TANDEM REPEATS. DROSOPHILA MELANOGASTER (FRUIT FLY). EUKARYOTA; METAZOA; ARTHROPODA; INSECTA; DIPTERA. (REL. 22, CREATED) (REL. 22, LAST SEQUENCE UPDATE) (REL. 31, LAST ANNOTATION UPDATE) PRT; 1002 AA. POLY-SER. NUCLEAR PROTEIN; DNA-BINDING; REPEAT. -!- PTM: THE N-TERMINAL IS BLOCKED TRANSCRIPTIONAL REGULATOR. STANDARD; 595 rsrkcvprkingsn 608 217 261 253 241 253 01-MAY-1992 (REL. YEMANUCLEIN-ALPHA. SEQUENCE FROM N.A. STRAIN=CANTON-S; MEDLINE; 92297435. YEMA OR YG4.5 01-MAY-1992 01-FEB-1995 YEMA DROME P25992; SEQUENCE VARIANT DOMAIN DOMAIN DOMAIN REPEAT REPEAT DOMAIN 유 ò

US-08-487-283A-1.rsp

Jan 28 12:17

ZEA MAYS (MAIZE). EUKARYOTA; PLANTA; EMBRYOPHYTA; ANGIOSPERMAE; MONOCOTYLEDONEAE; CYPERALES; GRAMINEAE.

SEQUENCE FROM N.A.
STRAIN-CV. INBRED LINE VA26;
BELANGER F.C., KRIZ A.L.;
PLANT PHYSIOL. 91:636-643(1989).

SEQUENCE OF 87-100.

01-AUG-1990 (REL. 15, 1AST SEQUENCE UPDATE) 01-OCT-1994 (REL. 30, LAST ANNOTATION UPDATE) GLOBULIN-1 S ALLELE PRECURSOR (GLB1-S) (7S-LIKE)

GLB1.

01-APR-1990 (REL. 14, CREATED)

573 AA.

STANDARD;

LT 8 GLB1_MAIZE

P15590;

291 kngecirdkve 301 |::|:|:|| 8 KSSKCVRQKVE 18

g E E 8 Š ö ö PROKARYOTA; FIRMICUTES; IRREGULAR ASPOROGENOUS RODS; CORYNEFORM GROUP 177 177 INVOLVED IN HYDROLYSIS OF THE SUBSTRATE. 264 Aa, 29057 MW; 81A56865 CRC32; Gaps Gaps X-RAY CRYSTALLOGRAPHY (2.0 ANGSTROMS), AND REVISIONS TO 184 AND 232 ROMAO M.J., TURK D., GOMIS-RUETH F.-X., HUBER R.; J. MOL. BIOL. 226:1111-1130(1992). -!- CATALYTIC ACTIVITY: N-CARBAMOYLSARCOSINE + H(2)0 = SARCOSINE + ALIEN T., SAMSON C., FERRI L., NAROD S., MORGAN K., SIMARD J.; GENOMICS 28:530-542(1995).
-!- SIMILARITY: BELONGS TO THE L34E FAMILY OF RIBOSOMAL PROTEINS. EMBL; L38941; G1008856; -.
RIBOSOMAL PROTEIN. 01-OCT-1993 (REL. 27, CREATED)
01-OCT-1993 (REL. 27, LAST SEQUENCE UPDATE)
01-FEB-1995 (REL. 31, LAST ANNOTATION UPDATE)
N-CARBAMOYLSARCOSINE AMIDASE (EC 3.5.1.59) (N-CARBAMOYLSARCOSINE ROMMENS J.M., DUROCHER F., MCARTHUR J., TONIN P., LEBLANC J.F., ö ö DICTYOSTELIUM DISCOIDEUM (SLIME MOLD). EUKARYOTA; PROTOZOA; SARCOMASTIGOPHORA; SARCODINA; RHIZOPODA; Score 57; DB 8; Length 116; Pred. No. 4.57e-01; 6; Mismatches 1; Indels 4; Indels Score 57; DB 2; Length 264; Pred. No. 4.57e-01; PRT; 537 AA.
P15269; P08126;
01-AUC-1988 (REL. 08, CREATED)
01-APR-1990 (REL. 14, LAST SEQUENCE UPDATE)
01-EFB-1994 (REL. 28, LAST ANNOTATION UPDATE)
SPORE COAT PROTEIN SP70 PRECURSOR (BEEJIN PROTEIN). -!- COFACTOR: ONE SULFATE ION PER SUBUNIT.
-!- PATHWAY: DEGRADATION OF CREATININE TO GLYCINE.
PIR; S28969; S28969.
PDB; INBA; 22-JUN-94.
HYDROLASE; 3D-STRUCTURE. SEQUENCE 116 AA; 13174 MW; 490F4AF1 CRC32; 5; Mismatches 264 AA. BY SIMILARITY. PRT; Query Match 40.4%; Best Local Similarity 41.7%; 40.48; -!- SUBUNIT: HOMOTETRAMER. 40.08; 5; Conservative AMIDOHYDROLASE) (CSHASE). 6; Conservative 171 gataagcvrhtveda 185 STANDARD; 0 6 GTKSSKCVRQKV 17 79 gsmcakcvrdri 90 Local Similarity MEDLINE; 92389321. CO(2) + NH(3). MEDLINE; 96039267 ARTHROBACTER SP CSH ARTSP ACT SITE SEQUENCE INIT MET Query Match Query Match Matches **fatches** RESULT
1D SP
1D SP
DT 011
DT 011
DE SP
GN CC
GN CC
OC EU RESULT g 8 £ 合 8

ö

Gaps ö COMER R.H., DATTA S., FIRTEL R.A.;
J. CELL BIOL. 103:1999-2015(1986).
EMBL; M26238; G167889; -.
PIR, B33485; B33485.
PIR, B25439; B25439.
DICTYNB, DD03009; COTB.
GLYCOPROTEIN; PHOSPHORYLATION; REPEAT; SPORULATION; SIGNAL. Score 56; DB 9; Length 537; Pred. No. 7.54e-01; 1; Indels 5.5 X 11 AA TANDEM REPEATS SPORE COAT PROTEIN SP70 MW; 5D59CBAC CRC32; PRESPORE MOTIF. PRESPORE MOTIF. 4; Mismatches 6 (INCOMPLETE) SER/THR-RICH, PRELIMINARY SEQUENCE OF 72-170 FROM N.A. POTENTIAL. FOSNAUGH K.L., LOOMIS W.F.; MOL. CELL. BIOL. 9:5215-5218(1989). 39.7%; Best Local Similarity 54.5%; 56650 EUMYCETOZOA; DICTYOSTELIA Conservative 20 537 250 248 200 211 222 233 244 248 248 263 263 371 537 AA; MEDLINE; 87057653. SEQUENCE FROM N.A. MEDLINE; 90097939. 266 9: REPEAT REPEAT CARBOHYD Query Match SEQUENCE SIGNAL DOMAIN DOMAIN REPEAT REPEAT REPEAT REPEAT REPEAT REPEAT REPEAT CHAIN Matches

O,

9

38.3%; Score 54; DB 2; Length 148; 63.6%; Pred. No. 2.01e+00; PIR, A32494, A32494. HYPOTHETICAL PROTEIN; TRANSPOSABLE ELEMENT. SEQUENCE 775 AA; 82355 MM; 9738B05A CRC32; 148 AA; 16534 MW; B5442F5E CRC32; 01-OCT-1994 (REL. 30, CREATED) 01-OCT-1994 (REL. 30, LAST SEQUENCE UPDATE) 01-FEB-1995 (REL. 31, LAST ANNOTATION UPDATE) SPORE COAT PROTEIN 2. 775 AA. 4; Mismatches 148 AA. -!- SUBCELLULAR LOCATION: SPORE OUTER COAT. ZHANG J., FITZ-JAMES P.C., ARONSON A.I.; J. BACTERIOL. 175:3757-3766(1993). GARRETT J.E., KNUTZON D.S., CARROLL D.; MOL. CELL. BIOL. 9:3018-3027(1989). EMBL; M26915; G214845; -. PRT; PRT; Best Local Similarity 46.78; Matches 7: Carr Query Match 39.7%; Best Local Similarity 41.2%; 450 idhlqnrrvrcvqemae 466 2 IDHQGTKSSKCVRQKVE 18 7; Conservative EMBL; L10116; G304149; -. PPIR; E47119; E47119. SUBTILIST; BG10500; COTZ. -!- SIMILARITY: TO COTY. STANDARD; 617 sntskcvssevegtp 631 STANDARD; :::|||| |||:: 7 TKSSKCVRQKVEGSS 21 STRAIN=168 / JH642; MEDLINE; 93285989. Best Local Similarity SEQUENCE FROM N.A. MEDLINE; 89384562. SEQUENCE FROM N.A. BACILLUS SUBTILIS LT 10 YTX1 XENLA SPORULATION. LT 11 COTZ BACSU SEQUENCE Query Match Q08312; $P143\overline{8}0;$ Matches RESULT RESULT В 유 ð Š ö MEDLINE, 89117617.
BORODIN A.M., DANILKOVICH A.V., CHERNOV I.I., AZHYKINA T.L.,
ROSTAPSHOV V.M., MONASTYRSKAYA G.S.;
BIOONG, KHIM. 14:1179-1182(1988).
--i- FUNCTION: DNA-DEPENDENT RNA POLYMERASE CATALYZES THE TRANSCRIPTION OF DNA INTO RNA USING THE FOUR RIBONUCLEOSIDE TRIPHOSPHATES AS -!- PTM: THREE PROTEIN-PROCESSING STEPS OCCUR IN THE FORMATION OF THE MATURE PROTEIN FROM THE PRIMARY TRANSLATION PRODUCT. -!- POLYMORPHISM: THE THREE MOST COMMONLY OCCURING GLB1 ALLELES HAVE Gaps CATALYTIC ACTIVITY: N NUCLEOSIDE TRIPHOSPHATE = N PYROPHOSPHATE SUBUNIT: THE EN2YME CONSISTS OF THE SIGMA CHAIN AND THE CORE ENZYME WHICH IS COMPOSED OF 2 ALPHA CHAINS, 1 BETA CHAIN, AND 1 01-NOV-1990 (REL. 16, CREATED) 01-MAR-1992 (REL. 21, LAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) DNA-DIRECTED RNA POLYMERASE BETA CHAIN (EC 2.7.7.6) (TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT). THE DESIGNATION L, I, AND S FOR LARGE, INTERMEDIATE, AND SMALL PROTEINS, RESPECTIVELY. BORODIN A.M., DANILKOVICH A.V., ALLIKWETS R.L., ROSTAPSHOV V.M., CHERNOV I.P., AZHIKINA T.L., MONASTYRSKAYA S., SVERDLOV D.; DOKL. BIOCHEM. 302:1261-1265(1988). PROKARYOTA; GRACILICUTES; SCOTOBACTERIA; AEROBIC RODS AND COCCI; BETA' CHAIN. -!- SIMILARITY: BELONGS TO THE RNA POLYMERASE BETA CHAIN FAMILY. ö Score 56; DB 4; Length 573; 2; Indels T -> N (IN REF. 2). I -> V (IN REF. 2). F -> S (IN REF. 2). 1357 AA; 151305 MW; BBF88A37 CRC32; Pred. No. 7.54e-01; OR 21 (POTENTIAL) 7E755E20 CRC32, 3; Mismatches PRT; 1357 AA EMBL; X15849; G45729; -. EMBL; M38319; G151547; -. TRANSCRIPTION; DNA-DIRECTED RNA POLYMERASE. GLOBULIN-1 S. POTENTIAL. [2] SEQUENCE OF 1036-1357 FROM N.A. 65029 MW; SIGNAL. 39.7%; 58.3%; EMBL; M24845; G168481; -. 7; Conservative STANDARD; 18 86 573 349 MAIZEDB; 30181; -. SEED STORAGE PROTEIN; 19 87 5° 849 345 673 AA; 32 hgghksgrcvrr 43 | | ||::|||: 4 HQGTKSSKCVRQ 15 HSSP; P02853; 1CAU. Best Local Similarity PSEUDOMONAS PUTIDA. [1] SEQUENCE FROM N.A. PSEUDOMONADACEAE. 1180 SUBSTRATES.

LT 9 RPOB PSEPU P19175;

CARBOHYD SEQUENCE

PROPEP SIGNAL CHAIN

222222EEEEEE

Query Match

datches

숌 ð RNA (N).

-:-+

CONFLICT

SEQUENCE

CONFLICT

ö ö PROKARYOTA; FIRMICUTES; ENDOSPORE-FORMING RODS AND COCCI; BACILLACEAE. EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; AMPHIBIA; ANURA Gaps Gaps -!- SUBUNIT: DISULFIDE CROSS-LINKED EITHER TO ITSELF OR TO COTY. ; ö Score 56; DB 8; Length 1357; Pred. No. 7.54e-01; Score 55; DB 11; Length 775; Pred. No. 1.24e+00; 5; Mismatches 3; Indels Indels 01-JAN-1990 (REL. 13, CREATED)
01-JAN-1990 (REL. 13, LAST SEQUENCE UPDATE)
01-APR-1990 (REL. 14, LAST ANNOTATION UPDATE)
TRANSPOSON TXI HYPOTHETICAL 82 KD PROTEIN (ORF 1).
XENOPUS LAEVIS (ARRICAN CLAMED FROG).

| ## CONSERVATIVE US-08-487-283A-LINSP CONSERVATIVE 14 |
|---|
| |

| LRRLET (IN RE) REF. 1)> DFKRGKERKF K -> QSVLSMGNL/ (IN REF. 1). GRC32; | ., . | | AA. | E) ATE) CHROMOSON | ; SECERNEN | DATA BANKS. (LRR) SEGMENT IS FOUND: 3. | AT. REPEATS. CRC32; | DB 11; Length 1039; 2.01e+00; atches 7; Indels | | AA. 5) NTE) |
|---|---|--|----------------------------|--|--|--|--|--|---|---|
| FEEIGNE -> D -> N (IN TSNVVRNESS AKRAFDGQFV FSYLHTMYI W; 8D658EBB | Score 54; DB 3; Lk Pred. No. 2.01e+00; 7; Mismatches 7; | : 531 | PRT; 1039 AA | QUSD'84; 01-NOV-1995 (REL. 32, IAST SEQUENCE UPDATE) 01-NOV-1995 (REL. 32, IAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, IAST ANNOTATION UPDATE) HYPOTHETICAL 118.2 KD PROTEIN F43C1.1 IN CHROMOSOME III | F43C1.1. GENORHABDITIS ELEGANS. EUKARYOTA; METAZOA; ACOELOMATES; NEMATODA; SECERNENTEA; RHABDITIDA SEQUENCE FROM N.A. | STRAIN=BRISTOL N2; JASSAL B.; SUBMITTED (DEC-1994) TO EMBL/GENBANK/DDBJ DATA BANKS SIMILARITY: THE REPEATED LEUCINE-RICH (LRR) SECM- MANY PROTEINS. NUMBER IN THIS PROTEIN: 3. | EMBL; 24693; G603526; WORMPEP; F43C1.1; CE01582. WORMPEP; F43C1.1; CE01582. DOMAIN 216 284 LEUCINE-RICH REPEAT. REPEAT 216 238 LRR 1. REPEAT 239 261 LRR 2. REPEAT 262 284 LRR 2. DOMAIN 669 903 PP2C-LIKE. SEQUENCE 1039 AA; 118182 MW; 877F95CB CRC32; | Score 54; Pred. No. 5; Mism | 697 20 | T 14 P04489; P04489; 13-AUG-1987 (REL. 05, CREATED) 11-AUG-1987 (REL. 05, LAST SEQUENCE UPDATE) 01-MAR-1992 (REL. 11, LAST ANNOTATION UPDATE) PRODBALE EARLY E4 11 KD PROTEIN. HUMAN ADENOVIRUS TYPE 5. |
| 182 188 200 200 366 375 808 818 818 AA; 93404 | 38.3%; ilarity 33.3%; Conservative | 511 ivdfhsrkaekcdkgnvlgks :: :: : 1 VIDHQGTKSSKCVRQKVEGSS | STANDARD; | (REL. 32, CREATED) (REL. 32, LAST SEG (REL. 33, LAST ANN L 118.2 KD PROTEIN | IIS ELEGANS. METAZOA; ACOELO OM N.A. | TOL N2; DEC-1994) TO EMITY: THE REPEAT OTEINS: NUMBER ITY: CONTAINS A | 7; G603526; 3C1.1; C601582. 21.1; C601582. 216 284 216 28 239 261 262 284 669 903 | 38.3%; ilarity 38.1%; Conservative | vagsrgmknkqcirqvrvent : : : : : VIDHQGTKSSKCVRQ-KVEGS | STANDAE REL. 05, REL. 21, X E4 11 F |
| FT CONFLICT FT CONFLICT FT CONFLICT FT CONFLICT FT CONFLICT FT CONFLICT | Query Match Best Local Similarity Matches 7; Conse | Db 511 ivdfhs: :: :: Qy 1 VIDHQG | RESULT 13 ID YR71 CAEEL | 01-3554; 01-NOV-1995 01-NOV-1995 01-FEB-1996 HYPOTHETICAL | GN F43C1.1. 0S CAENORHABDITIS ELEGANS 0C EUKARYOTA; METAZOA; AC N [1] RP SEQUENCE FROM N.A. | STRAIN-BRISTOL N2; ASSAL B.; SUBMITTED (DEC-199 -!- SIMILARITY: TH MANY PROTEINS. | EMBL; 24693 WORMPEP; F47 HYPOTHETICAL DOMAIN REPEAT REPEAT REPEAT REPEAT SEQUENCE 1 | Query Match Best Local Similarity Matches 8; Conse | 677 vsgsrgn : 1 VIDHQG1 | RESULT 14 AC E411 ADEO5 STANDAF AC PO4489; DT 13-AUG-1987 (REL. 05, DT 13-AUG-1987 (REL. 05, DT 01-MAR-1992 (REL. 21, DF PROBABLE EARLY E4 11 K OS HUMAN ADEMOVIEUS TYPE |

```
;
0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         P03241;
21-JUL-1986 (REL. 01, CREATED)
21-JUL-1986 (REL. 01, LAST SEQUENCE UPDATE)
01-APR-1990 (REL. 14, LAST ANNOTATION UPDATE)
HUMAN ADENOVIRUS TYPE 2.
VIRIDAE; DS-DNA NONENVELOPED VIRUSES; ADENOVIRIDAE; MASTADENOVIRUSES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        .;
0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
                                                                                                        MEDLINE; 83164198.
SARNOW P., HEARING P., ANDERSON C.W., REICH N., LEVINE A.J.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           [1]
SEQUENCE FROM N.A.
MEDLINE; 82059444.
MEDLINE; 82059444.
NUCLEIC ACIDS RES. 9;4023-4042(1981).
EMBL; J01917; G209839; -.
PIR; A03807; Q4ADE2.
EXALY PROTEIN.
US-08-487-283A-1.rsp
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match 37.6%; Score 53; DB 3; Length 116; Best Local Similarity 60.0%; Pred. No. 3.25e+00; Matches 6; Conservative 3; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 37.6%; Score 53; DB 3; Length 116; Best Local Similarity 60.0%; Pred. No. 3.25e+00; Matches 6; Conservative 3; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       SEQUENCE 116 AA; 13255 MW; 950D6981 CRC32;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SEQUENCE 116 AA; 13298 MW; 66EA9B5C CRC32;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PRT; 116 AA.
                                                                                                                                                                                                                 [2]
COMPLETE GENOME.
MEDLINE; 92087470.
CRROBOCZEK J., BIEBER F., JACROT B.;
CRROBOCZEK J., BIEBER F., JACROT B.;
EMBL; M73260; -; NOT_ANNOTATED_CDS.
EMBL; X02998, G58502; -.
PIR; B03807; QAADES.
                                                                                                                                                                                     I. MOL. BIOL. 162:565-583(1982).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            6; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    :|:| |||||:
11 KCVRQKVEGS 20
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              3 rclrlkvega 12
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       JLT 15
E411_ADE02
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT
1D E41
AC 2013
DT 21-2
DT 21-2
DT 21-2
DT 01-2
DE PRC
OS HUR
RN [1]
RN [1]
RN HEI
RA HEI
RA HEI
RA HEI
RA HEI
RA KEI
RA K
                                                                                                               g
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ð
```

Search completed: Wed Jan 28 12:11:02 1998 Job time : 10 secs.

3 rclrlkvega 12 :|:| ||||: 11 KCVRQKVEGS 20

임 ð

US-08-487-283A-1.rag Jan 28 12:16 Œ

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 Unive.aity of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

protein - protein database search, using Smith-Waterman algorithm MPsrch_pp

Wed Jan 28 12:09:44 1998; MasPar time 2.31 Seconds 111.852 Million cell updates/sec Run on:

Tabular output not generated.

>US-08-487-283A-1 (1-21) from US08487283A.pep Description: Title:

1 VIDHQGTKSSKCVRQKVEGSS 21 Perfect Score: Sequence:

PAM 150 Gap 15 Scoring table:

101610 seqs, 12294212 residues Searched:

Minimum Match 0% Listing first 45 summaries Post-processing:

Database:

i.parti 2:part2 3:part3 4:part4 5:part5 6:part6 7:part7 8:part8 9:part9 10:part10 11:part11 12:part12 13:part13 14:part14 15:part15 16:part16 17:part17 18:part18 19:part19 20:part20 21:part21 a-geneseq28

Mean 18.854; Variance 56.108; scale 0.336 Statistics: Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| <u>.</u> | 6 | 6 | = | | _ | = | _ | _ | - | = | = |
|--------------------------|-----------------------|---------------------|---------|-------------------------|-----------------------|-----------------------|----------------------|-----------------------|-----------------------|-----------------------|-------------|
| Pred. No. | 1.23e-09 | | m | 4.47e+01 | 4.47e+01 | 5.69e+0 | 5.69e+01 | 5.69e+0 | 5.69e+0 | 5.69e+01 | 5.69e+0] |
| Description | Pro-C5 polypeptide KS | Pro-C5 polypeptide. | CSHase. | . P. gingivalis porphyp | Hepatitis GB virus (H | Mouse epithelin precu | Tobacco mosaic virus | Type B human platelet | Alpha type PDGF recep | Platelet derived grow | Rat petrin. |
| 11 | | R77604 | R22271 | R96029 | R94347 | R14327 | R88124 | R26206 | R06910 | R08267 | W04326 |
| 80 | 15 | 15 | 4 | 11 | 16 | m | 15 | S | 7 | 7 | 19 |
| Query Match Length DB | 21 | 1676 | 264 | 1732 | 3163 | 589 | 652 | 1009 | 1089 | 1089 | 1196 |
| Query Match | 100.0 | 100.0 | 39.0 | 38.3 | 38.3 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 | 37.6 |
| Score | 141 | 141 | 22 | 24 | 24 | 53 | 53 | 23 | 53 | 53 | 23 |
| Result No. | 1 | 7 | e | 4 | 'n | ۍ | 7 | 00 | o | 10 | Π |

US-08-487-283A-1.rag Jan 28 12:16

N

| 12 | 52 | 36.9 | 354 | 9 | R33439 | Ornithine cyclodeamin | 7.25e+01 |
|----|----|------|------|----|--------|-----------------------|------------|
| 13 | 51 | | 589 | ٣ | R14325 | Rat epithelin precurs | 9.21e+01 |
| 14 | 51 | 36.2 | 914 | ٣ | R15785 | | 9.21e+01 |
| 15 | 51 | | 926 | m | R15784 | B.thuringiensis toxin | 9.21e+01 |
| 16 | 51 | | 986 | 6 | R25141 | JAK2. | 9.21e+01 |
| 17 | 51 | | 1100 | m | R15783 | B.thuringiensis toxin | 9.21e+01 |
| 18 | 51 | 36.2 | 1129 | 13 | R70830 | Murine JAK2 kinase. | 9.21e+01 |
| 19 | 21 | | 1144 | 12 | R88122 | Tobacco mosaic virus | 9.21e+01 |
| 20 | 51 | | 1144 | 15 | R88123 | Tobacco mosaic virus | 9.21e+01 |
| 21 | 21 | | 1588 | 6 | R46605 | Malarial PfEMP3 epito | 9.21e+01 |
| 22 | 51 | 36.2 | 1663 | 6 | R46608 | Plasmodium falciparum | 9.21e+01 |
| 23 | 20 | | 1047 | 20 | W01535 | Cellular homologue of | 1.17e+02 |
| 24 | 49 | _ | 429 | 10 | R50036 | Hantaian virus Nucleo | 1.48e + 02 |
| 25 | 49 | | 429 | 21 | W15472 | Hantaan virus nucleoc | 1.48e + 02 |
| 56 | 49 | 34.8 | 429 | 20 | W11872 | Hantavirus nuclear pr | 1.48e+02 |
| 27 | 49 | | 430 | 50 | W11874 | Hantavirus nuclear pr | 1.48e+02 |
| 88 | 49 | | 430 | 20 | W11873 | Hantavirus nuclear pr | 1.48e+02 |
| 59 | 49 | 34.8 | 487 | m | R13794 | normone | 1.48e+02 |
| 30 | 49 | | 694 | - | R04107 | DNA-binding protein G | 1.48e+02 |
| 31 | 49 | 34.8 | 1323 | 10 | R55248 | | 1.48e + 02 |
| 32 | 49 | | 1336 | 12 | R66041 | Human N-methyl-D-aspa | 1.48e + 02 |
| 33 | 49 | • | 3080 | - | P93285 | Sequence of clone HIV | 1.48e+02 |
| 34 | 48 | 34.0 | 399 | 18 | W04218 | Human urinary bladder | 1.87e+02 |
| 32 | 48 | | 925 | 14 | R79148 | Human insulin recepto | 1.87e+02 |
| 36 | 48 | 34.0 | 1299 | 15 | R86304 | Drosophila patched pr | |
| 37 | 47 | | 122 | 16 | R81443 | Hepatitis GB virus (H | 2.36e+02 |
| 38 | 47 | | 458 | m | R15149 | CD4 coordinate system | 2.36e+02 |
| 39 | 47 | | 498 | ٣ | R12255 | HIV-1 strain OYI GAG | |
| 40 | 47 | | 792 | 16 | R85198 | Avenacinase - a sapon | 2.36e+02 |
| 41 | 47 | | 793 | 16 | R85200 | Avenacinase-like prot | 2.36e+02 |
| 42 | 47 | | 793 | 16 | R85199 | a | 2.36e+02 |
| 43 | 47 | 33.3 | 2873 | 11 | R87559 | rus c | 2.36e+02 |
| 44 | 47 | | 2873 | 11 | 79 | lypr | 2.36e+02 |
| 45 | 46 | 32.6 | 1232 | 19 | R98217 | Neuronal apoptosis in | 2.97e+02 |

ALI GNMENTS

Pro-C5 polypeptide KSSKC epitope. Complement C5; haemolysis; kidney; glomerulonephritis; monoclonal antibody; antiinflammatory; antibody engineering; R77605 standard; Protein; 21 AA. 02-APR-1996 (first entry) R77605; RESULT

humanised antibody; KSSKC epitope. 02-MAY-1994; US-236208. (ALEX-) ALEXION PHARM INC. 01-MAY-1995; U05688 Homo sapiens. WO9529697-A1. 09-NOV-1995

Treating glomerulonephritis with antibody against complement C5 Evans MJ, Matis L, Mueller EE, Nye SH, Rollins S; Rother RP, Springhorn J P, Squinto SP, Thomas TC; - to inhibit complement induced cell lysis Example 13; Page 81; 181pp; English. Rother RP, Springhor Wang Y, Wilkins JA; WPI; 95-392923/50. component

The cDNA sequency of the complement C5 gene transcript predicts a secreted pro-C5 precursor of 1676 amino acids (R77604). C5 is a beta-globulin heterodimer thought to play a role in the pathogenesis of glomerulonephritis (GN). Cleavage of the C5 alpha-chain by a convertase enzyme generates anaphylatoxic C5a. Monoclonal

ť

នួននន

셤

ð

m

Jan 28 12:16

ö Gaps and humanised recombinant antibodies that recognise the alpha-chain KSSKC epitope (R77605) block C5a generation, thereby reducing glomerular inflammation and kidney dysfunction associated with GN. ; 0 Score 141; DB 15; Length 1676; Pred. No. 1.23e-09; 0; Indels 0; Mismatches Query Match 100.0%; Best Local Similarity 100.0%; 21; Conservative 1676 AA:

efficient expression in E.coli
Claim 9; Page 9 + 7; 12pp; German.
The sequence encoding CSHase is useful in assay of creatinine
(for diagnosis of kidney disease). It can now be prepd. more
simply than by known methods which involve culture of Arthrobacter N-PSDB; 022713.
Recombinant DNA encoding N-carbamoyl-sarcosine-amidohydrolase useful in clinical assay of creatinine, and vectors providing N-carbamoyl-sarcosine amidohydrolase; CSH; assay; diagnosis; J. 3 R22271 standard; Protein; 264 AA. (BOEF) BOEHRINGER MANNHEIM GMBH. 30-JUL-1992 (first entry) Burtscher H, Schumacher G; WPI; 92-098378/13. Arthrohacter sp. DSM 2563 20-SEP-1990; DE-029844. 19-SEP-1991; 115974.

Score 55; DB 4; Length 264; Pred. No. 3.50e+01; 39.08; Best Local Similarity 46.2%; 264 AA; on complex media.

ö Gaps ö 3; Indels 4; Mismatches 6; Conservative

171 gataagcvrhtve 183 |: :: |||: || 6 GTKSSKCVRQKVE 18

Porphypain; haemagglutinin; periodontal disease; vaccine; antibody. Porphyromonas gingivalis strain W12. Location/Qualifiers 04-SEP-1996 (first entry) P. gingivalis porphypain. 688..708

"Pro-Asn repeat region type 1" 887..952 "Pro-Asn repeat region type 2" 946..967

985..1006

/note= "Pro-As WO9617936-A2.

Region /note= /note=

Region Region Region

Region

13-JUN-1996.

Tumwasorn S;

(UYFL) Han N,

S

US-08-487-283A-1 rag

Sequence Query Match

Matches

쇰 ð

Gaps Non-A, non-B, non-C, non-D, non-E Hepatitis virus reagents - useful for diagnosis and therapy of hepatitis GB virus
Example 9; Pages 401-414; 661pp; English.
Double strandad hepatitis GB virus (HGBV) DNA obtd. from HGBV found to be unique HGBV sequences. The clones were then used to assemble the sequences T00129/30 (GB contig A and B) which encode the proteins R94345-47 (the 3 possible coding strand reading frames) and R82072, respectively. Reagents which comprise the HGBV DNA, or its protein prods. can be used for the diagnosis, therapy or in a vaccine to prevent HGBV infection. infected tamarin plasma, using standard procedures, was used to prepare a lambda phage HGBV cDNA library. Clones were rescued from the lambda phage, searched against a sequence database and 4; Indels 0; Buijk SL, Dawson GJ, Desai SM, Erker JC, Leary TP; Muerhoff AS, Mushahwar IK, Pilot-Matias TJ, Schlauder GG; Score 54; DB 16; Length 3163; Pred. No. 4.47e+01; ET; growth regulation; inhibition; stimulation. 5; Mismatches Dawson GJ, Desai SM, Erker JC,

/label= EP /note= "claim 27, page 56"

US-196030

4-FEB-1994;

4-FEB-1995; U02118.

.7-AUG-1995.

13-MAY-1994; US-242654 29-JUL-1994; US-283314

Peptide

/label: EP

Peptide

ö

US-08-487-283A-1.rag

/note= "claim 28, page 56"

Jan 28 12.16

03-APR-1991; U02321.

Sequence Query Match

Aatches

a ð

17-0CT-1991. W09115510-A

Alpha type PDGF receptor deduced from TR4 cDNA clone. Platelet derived growth factor; T11. PDGF activity in sample Disclosure; Page 90; 109pp; English. Location/Qualifiers LT 9 R06910 standard; protein; 1089 AA. R26206 standard; Protein; 1009 AA. 381 vddhhgstggqtvrctaegtp 401 21 (CORT-) COR THERAPEUTICS INC. Query Match 37.6%; Best Local Similarity 38.1%; R26206; 09-FEB-1993 (first entry) 1 VIDHQCTKSSKCVRQKVEGSS 16-JAN-1991 (first entry) 8; Conservative 24..1009 31-JAN-1991; US-650793. 'label= Signal_peptide /label= Mature_PDGF-A 28-JAN-1992; U00730 1009 AA; bind a PDGF Ligand. WPI; 92-299970/36. N-PSDB; Q27451. Homo sapiens. W09213867-A. 20-AUG-1992. Sequence Protein Peptide R06910; Matches RESULT a ₩ E E E E ò ö ö ET-1 is a bifunctional growth regulator, capable of stimulating the growth of some cell types while inhibiting the growth of others. ET-2 is functionally similar to ET-1 w.r.t. growth inhibitory bioactivity. In contrast, however, ET-2 is apparently not capable of eliciting the growth stimulatory activity characteristic of ET-1 and, generating transgenic Solanaceous plants resistant to Tobacco Mosaic Gaps Gaps protein can be used to generate transgenic plants, esp. Solanaceae, The Nicotiana glutinosa N gene truncated protein (R88124) mediates resistance to tobacco mosaic virus (TWV). A cDNA clone (T09342) New cysteine-rich growth modulating proteins, epithelins - useful as inhibitors of neoplastic cell growth and to promote wound Plant virus resistance gene N sequences from tobacco - useful for coding for the protein was obtd. from a N. glutinosa leaf cDNA ö ö library by transposon tagging. DNA sequences encoding the Score 53; DB 15; Length 652; Pred. No. 5.69e+01; Tobacco mosaic virus resistance; TMV; N gene; Solanaceae; 6; Indels Length 589; 2; Indels Tobacco mosaic virus resistance N gene truncated protein. in fact, antagonises this ET-1 activity. See also Q14338-40, Q14952-53, R14328-9 and R15315-20. crop improvement; transgenic plant; crop improvement. Score 53; DB 3; Lk Pred. No. 5.69e+01; 7; Mismatches 5; Mismatches Claim 28; Page 75-79; 98pp; English. Disclosure, Fig 23; 97pp; English. R88124 standard; Protein; 652 AA. (BRIM) BRISTOL-MYERS SQUIB. healing and treat psoriasis 35.7%; 37.6%; 31.3%; 28-MAR-1996 (first entry) 5; Conservative 28-DEC-1995. 16-UNN-1995, U07754. 17-UNN-1994, US-261663. (REGC) UNIV CALIFORNIA. (USDA) US SEC OF AGRIC. Baker BJ, Whitham SA; Conservative 03-APR-1990; US-504508. 13-MAR-1991; US-083796. 560 hcsargtkclrkki 573 4 HQGTKSSKCVRQKV 17 Shoyab M, Plowman GD; Nicotiana glutinosa. W09535024-Al Query Match Best Local Similarity Best Local Similarity WPI; 91-325168/44. N-PSDB; Q14340. WPI; 96-058144/06. resistant to TMV. Sequence 652 AA;

R88124;

RESULT

US-08-487-283 A-1 rag Jan 28 12:16

156 dnrdktdadcirqivd 171

a

two transmembrane glycoproteins, each of which is about 180 kD. This receptor has three major regions. The first is a transmembrane region, which spans the membrane once, separating the regions of the Gaps which possesses a tyrosine kinase activity. This tyrosine kinase domain is notable in having an insert of approx. 100 amino acids, as compared with most other receptor tyrosine kinase domains which are contiguous or have shorter insert sequences. Fragments of this sequence between 8 and 400 amino acids comprising one or more PDGF growth factor (PDGF) receptor (PDGF-R). This receptor is typically found on cells of mesenchymal origin. It acts while in the form of receptor exterior to the cell from those interior to the cell. The second region is an extracellular region which contains the domains igand binding region from the extracellular domain may be used to Type B human platelet-derived growth factor receptor. PDGF; PDGF-R; mesenchyme; tyrosine kinase; ligand binding region. which bind the PDGF. The third region is an intracellular region The sequence given is one allele of type A human platelet-derived Platelet derived growth factor receptor (PDGF-R) poly:peptide(s) - useful as therapeutic and diagnostic agents e.g. for assaying ; 0 Escobedo JA, Fretto LJ, Giese NA, Tomlinson JE, Williams LT; Score 53; DB 5; Length 1009; Pred. No. 5.69e+01; 7; Indels 6; Mismatches 1:: : |:| |: 3 DHQGTKSSKCVRQKVE 18 8

ö

N-PSDB; T09342.

Virus

Matches

Ó

susceptible to genetic engineering.
Claim 7: Fig 3; 64pp; English.
The TR4 clone is the largest Colon clone related to the Tl1 genomic clone, isolated from a inbrary prepd. from human thymus DNA. The T4 cDNA clone was isolated from a M426 human embryo fibroblast cDNA library. The coding region can be introduced into the pSV2 plasmic region comprises a juxtamembrane region, a tyrosine kinase region split into TKI and TK2 by a hydrophilic interkinase region and a hydrophilic C-terminal trail. The TK region includes the consensus ATP binding sequence (G-X-G-X-K-C...K) and a tyrosine autophosphorylation site homologous to that of pp60(v-src). type beta). The polypeptide has a calculated molecular mass of 120 kinase receptor. The extracellular region comprises a hydrophobic kD and has all the characteristics of a membrane spanning tyrosine for transforming cells to express novel protein receptor and also gpt vector with a simian sarcoma virus LTR as a promoter and expressed in a host. The resulting protein is a novel PDGF receptor designated type alpha (the known receptor is designated Type alpha platelet-derived growth factor teceptor gene - useful homology with the PDGF-R/CSF1-R subfamily. Ten Cys residues are family and eight potential N-linked glycosylation sites are also present. A hydrophobic segment spans the membrane and the cytosignal peptide and a ligand binding domain which has structural spaced at the same positions as in other receptors of the sub-/label=tyrosine autophosphorylation site Location/Qualifiers Matsui T, Aaronson SA, Pierce JH; /label-ligand binding domain Domain 525..548 /label-transmembrane region Domain 549..599 /label=juxtamembrane domain (USDC) US SEC OF COMMERCE. Tabel=N-glycos site /label=N-glycos_site Modified-site 353..355 /label=N-glycos_site Modified-site 359..361 /label=N-glycos_site Modified-site 468..470 /label=N-glycos_site 4odified-site 103..105 600..627 /label=N-glycos_site Modified-site 179..181 24..524 /label=ATP binding site /label=N-glycos_site 09-FEB-1989; US-308282 /label=signal peptide 'label=N-glycos_site 08-FEB-1990; U00617. Modified-site 849 WPI; 90-290306/38. Modified-site Andified-site Modified-site Modified-site N-PSDB; 005989. Homo sapiens. Binding-site W09010013-A. 07-SEP-1990. Sequence Omain)omain

ö Gaps ö Score 53; DB 2; Length 1089; Pred. No. 5.69e+01; 7; Indels US-08-487-283A-1.rag 6; Mismatches 421 vidhhgstggqtvrctaegtp 441 1 VIDHQGTKSSKCVRQKVEGSS 21 8; Conservative Best Local Similarity :: :: :: Jan 28 12:16 Query Match Matches අ à

R08267 standard; protein; 1089 AA.

RESULT

ö utility in dection of PDGF agonist and antagonist analogues, binding Gaps DNA encoding platelet-derived growth factor - used to transform ;; Gene product may be expressed from a transformed cell. It has AA, AB and BB isoforms. PDGF agonists may be used to enhance healing, and antagonists may be used to block the effects of eg. in treatment of atherosclerosis or fibrotic diseases. cells for culturing to detect PDG agonists and antagonists 37.6%; Score 53; DB 2; Length 1089; 38.1%; Pred. No. 5.69e+01; 7; Indels Platelet derived growth factor (PDGF) receptor protein. Atherosclerosis; fibrotic diseases. 6; Mismatches 421 vddhhgstggqtvrctaegtp 441 Claim 1; Fig 1; 30pp; English. 07-MAR-1991 (first entry) Best Local Similarity 38.1%; 8; Conservative (ZYMO-) ZYMOGENETICS INC. 22-MAY-1989; US-355018. Kelly JD, Murray MJ; WPI; 90-375992/50. N-PSDB; Q06869. 21-MAY-1990; U02849. Homo sapiens. W09014425-A. 29-NOV-1990. Sequence Query Match Matches 쇰

ð /F 11 W04326 standard; Protein; 1196 AA.

central nervous system; myelin; protein phosphatase 2C; stroke; Petrin; neurite outgrowth associated protein; CNS; Location/Qualifiers 16-JAN-1997 (first entry) Misc difference 129 neurodegeneration. Rat petrin. Rattus sp.

/note= "corresponds to stop codon in DNA sequence" Misc_difference 219 /note= "corresponds to stop codon in DNA sequence" "corresponds to stop codon in DNA sequence" Misc difference 192 Misc difference 205 /note=

/note= "corresponds to stop codon in DNA sequence" Misc difference 234 /note= "corresponds to stop codon in DNA sequence" Misc_difference 225

US-08-487-283A-1 rag

(MOUN) MOUNT SINAI HOSPITAL CORP

13-APR-1995; US-421701

US-08-487-283A-1.rag

/note= "corresponds to stop codon in DNA sequence" 'note= "corresponds to stop codon in DNA sequence" /note= "corresponds to stop codon in DNA sequence"

difference 269 difference 243

Aisc difference 285 Misc difference 312 Misc difference 319

/note= "corresponds to stop codon in DNA sequence"

corresponds to stop codon in DNA sequence"

/note= "corresponds to stop codon in DNA sequence" /note= "corresponds to stop codon in DNA sequence" Misc_difference 378 /note= "corresponds to stop codon in DNA sequence"

Misc difference 344 Aisc difference 358

"corresponds to stop codon in DNA sequence

'note= "corresponds to stop codon in DNA sequence" 'note= "corresponds to stop codon in DNA sequence"

12-APR-1996; CA0214.

409632476-A1. .7-0CT-1996.

'note= "corresponds to stop coodn in DNA sequence" 'note= "corresponds to stop codon in DNA sequence"

'note= "corresponds to stop coodn in DNA sequence" /note= "corresponds to stop coden in DNA sequence"

Misc difference 1017 Misc difference 1054 Misc difference 1127 Misc difference 1147 Misc_difference 1178

/note≂ "corresponds to stop codon in DNA sequence'

'note= "corresponds to stop codon in DNA sequence

Misc difference 924 disc difference 934

ö ö protein phosphatase 2C family, and is expressed in neurons in brain tissue, partic. in the Purkinje cells of the cerebellum. Petrin, and antibodies raised against it, can be used to modulate neurite Gaps Gaps Assay for substance that modulates response of neuronal cells - and mu-crystallins; drug targetting; nervous acting drugs; CNS; neural; antiepileptic agents; antianxiety agents; hallucinogens; sedatives; CNS depressants; CNS stimulators; tranquilisers; muscle relaxants; involving nerve damage resulting from traumatic injury, stroke or neurite growth associated protein, Petrin, useful in conditions Rat petrin (M04326) is a protein involved in modulating neurite growth inhibition. The amino sequence was deduced from a cDNA clone (T38484) derived from an adult rat brain cDNA library; no New mu-crystalline proteins - having ornithine cyclo-deaminase ; 0 coding sequence was indicated. Petrin is a new member of the ö neuronal; neurotransmitter agents; neuromuscular agents; NMJ; This sequence represents ornithine cyclodeaminase (OCD) from Agrobacterium Ti plasmid pTiC58. It shows approximately 30% Score 53; DB 19; Length 1196; Pred. No. 5.69e+01; neuromuscular junctions; memory agents; Alzheimers disease; homology with the kangaroo eye lens protein mu-crystallin. antispasmodics; analgesics; anesthetics; anticonvulsants; activity, used in diagnosis and treatment of disorders in Score 52; DB 6; Length 354; Pred. No. 7.25e+01; 4; Indels Indels Ornithine cyclodeaminase C58 from Ti plasmid pTiC58 ä 1; Mismatches Mismatches Lozano A, Roach A, Roder J; 01-Jan-1993. 28-FEB-1992; 844304. 28-FEB-1992; US-844304. (USSH) US DEPT HEALTH & HUMAN SERVICE. Claim 9; Page 57-61; 119pp; English. Disclosure; Page 34; 60pp; English. Jr 12 R33439 standard; Protein; 354 AA. growth and axonal regeneration. 37.6%; CNS degenerative disorders Query Match 36.9%; Best Local Similarity 63.6%; 06-JUL-1993 (first entry) 8; Conservative 7; Conservative Agrobacterium tumefaciens 900 vpshsgtksphcv 912 1 VIDHQGTKSSKCV 13 ornithine metabolism 312 ryvrdrvegss 322 Best Local Similarity WPI; 96-477127/47. WPI; 93-093573/11. 354 AA; N-PSDB; T38484 US7844304-A. hypnotics. Sequence Sequence Query Match Kim R, Matches Matches RESULT RESULLT RESULLT RESULLT RESULLT RESULLT RESULLT RESULLT RESULT RE g ð 용

/note= "corresponds to stop codon in DNA sequence" Misc difference 555

/note= "corresponds to stop coodn in DNA sequence" Misc_difference 593

/note= "corresponds to stop codon in DNA sequence" Misc_difference 602

/note= "corresponds to stop codon in DNA sequence" Misc_difference 609 'note= "corresponds to stop codon in DNA sequence' /note= "corresponds to stop codon in DNA sequence"

Misc difference 621 Misc difference 724

/note= "corresponds to stop codon in DNA sequence" /note= "corresponds to stop codon in DNA sequence" /note= "corresponds to stop codon in DNA sequence' /note= "corresponds to stop codon in DNA sequence' Misc_difference 841

Misc difference 736 Misc_difference 739 Misc difference 786

/note= "corresponds to stop codon in DNA sequence" Misc_difference 494

/note= "corresponds to stop codon in DNA sequence"

/note= "corresponds to stop codon in DNA sequence"

'note= "corresponds to stop codon in DNA sequence"

Misc difference 386

Misc difference 465 Misc difference 473 Misc difference 473

11 KCVRQKVEGSS 21

Jan 28 12:16

9

Jan 28 12:16

R15785 standard; Protein; 914 AA.

RESULT

US-08-487-283A-1.rag

```
ET-1 is a bifunctional growth regulator, capable of stimulating the growth of some cell types while inhibiting the growth of others. ET-2 is functionally similar to ET-1 w.r.t. growth inhibitory bloactivity. In contrast, however, ET-2 is apparently not capable of
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     eliciting the growth stimulatory activity characteristic of ET-1 and,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 New cysteine-rich growth modulating proteins, epithelins - useful as inhibitors of neoplastic cell growth and to promote wound
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 51; DB 3; Length 589;
Pred. No. 9.21e+01;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              See also Q14338-40, Q14952-53, R14328-9 and R15315-20.
                                                  ET; growth regulation; inhibition; stimulation.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  in fact, antagonises this ET-1 activity.
                                                                          Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          healing and treat psoriasis
Disclosure; Fig 18; 97pp; English.
R14325 standard; Protein; 589 AA.
                                                                                        Protein
/label= precutsor
/note= "claim 11, page 54"
280..335
                                                                                                                                                                                                                                                                                                                                                                                                                                                               (BRIM ) BRISTOL-MYERS SQUIB.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   36.2%;
                                                                                                                                                   /note= "claim 12, page 54"
Protein 205..261
                         (first entry)
                                                                                                                                                                                        /note= "claim 13, page 54"
                                                                                                                                                                                                                              /note= "claim 14, page 54"
                                                                                                                                                                                                                                                                     page 54"
                                                                                                                                                                                                                                                                                             /label= EP
/note= "claim 16, page 54"
                                                                                                                                                                                                                                                                                                                                               'note= "claim 17, page 54"
                                                                                                                                                                                                                                                                                                                                                                                     'note= "claim 18, page 55"
                                                                                                                                                                                                                                           123..179
                                                                                                                                                                                                                                                                                  362..416
                                        Rat epithelin precursor.
                                                                                                                                                                                                       59..114
                                                                                                                                                                                                                                                                                                                                                                                                                                    03-APR-1990; US-504508.
13-MAR-1991; US-083796.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Shoyab M, Plowman GD;
                                                                                                                                                                                                                                                                                                                                                                                                                         13-APR-1991; U02321.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WPI; 91-325168/44.
                                                                                                                                                                                                                                                                     'note= "claim 15,
                                                               Rattus rattus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     N-PSDB; Q14338
                       17-JAN-1992
                                                                                                                                     /label= EP-1
                                                                                                                                                                                                                                                                                                                                                                                                               7-0CT-1991.
                                                                                                                                                                              /label= EP-2
                                                                                                                                                                                                                                                                                                                                                                                                 409115510-A.
                                                                                                                                                                                                                   /label= EP
                                                                                                                                                                                                                                                         /label= EP
                                                                                                                                                                                                                                                                                                                                  /label= EP
                                                                                                                                                                                                                                                                                                                                                                         'label= EP
                                                                                      Protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
                                                                                                                                                                                                        Peptide
                                                                                                                                                                                                                                              eptide
                                                                                                                                                                                                                                                                                  eptide
                                                                                                                                                                                                                                                                                                                       eptide
                                                                                                                                                                                                                                                                                                                                                             eptide
              R14325;
```

Gaps carboxyl terminus coding region of B.thuringiensis var. tenebriosis carboxyl terminus coding region of B.thuringiensis var. tenebriosis (Btt) toxin. DNA encoding the gp64 viral membrane protein of AcNPV Extending host range or toxicity of insecticidal proteins - using three B.thuringiensis toxin/AcMPV gp64 fusion protein. chimeric; fusion protein; insecticide; AcMPV; Lepidoptera larvae; midgut targetting; bacterial endotoxin; pFX7. Extending host range or toxicity of insecticidal proteins - using chimeric; fusion protein; insecticide; AcNPV; Lepidoptera larvae; A polylinker was inserted into the XmnI restriction site at the A polylinker was inserted into the XmnI restriction site at the ; 0 different $\mbox{BtL/gp64}$ gene fusions that were constructed and its deduced amino acid sequence is given here. polylinker. The gp64 gene sequences act as midgut targetting was operably linked to the Btt toxin coding sequence via the signals for bacterial endotoxins. Plasmid pFAC13 was one of Score 51; DB 3; Length 914; Pred. No. 9.21e+01; 1; Indels protein capable of binding to gut epithelium of insects protein capable of binding to gut epithelium of insects Claim 55; Fig 17; 61pp; English. Autographa californica Nuclear Polyhedrosis Virus. W09117254-A. Autographa californica Nuclear Polyhedrosis Virus. B.thuringiensis toxin/AcNPV gp64 fusion protein. midgut targetting; bacterial endotoxin; pFAC13. 5; Mismatches Bacillus thuringiensis var. tenebriosis. Bacillus thuringiensis var. tenebriosis. R15784 standard; Protein; 956 AA. 14-NOV-1991. 02-MAY-1991; U03008. 03-MAY-1990; U5-518575. (REGC) UNIV OF CALIFORNIA. Sivasubramanian N, Federici A; Claim 55; Fig 18; 61pp; English. Sivasubramanian N, Federici A; See also Q14806 and Q14807. Sequence 914 AA; Query Match 36.2%; Best Local Similarity 45.5%; (REGC) UNIV OF CALIFORNIA. 10-FEB-1992 (first entry) 10-FEB-1992 (first entry) 5; Conservative 03-MAY-1990; US-518575. 665 kfnrcikrkve 675 | ::|:::||| 8 KSSKCVRQKVE 18 02-MAY-1991; U03008. WPI; 91-353775/48. WPI; 91-353775/48. N-PSDB; Q14807. N-PSDB; Q14808. 14-NOV-1991. W09117254-A. Matches DP à

ö

Gaps

ö

2; Indels

5; Mismatches

6; Conservative

쇰 ð

ö

US-08-487-283A | rag

(Btt) toxin. DNA encoding the gp64 viral membrane protein of AcNPW was operably linked to the Btt toxin coding sequence via the polylinker. The gp64 gene sequences act as midgut targetting singlals for bacterial endotoxins. Plasmid pFX7 was one of three different Btt/gp64 gene fusions that were constructed and its deduced amino acid sequence is given here. See also Q14806 and Q14808. Sequence

88888888

Gaps <u>`</u> Query Match 36.2%; Score 51; DB 3; Length 956; Best Local Similarity 45.5%; Pred. No. 9.21e+01; Matches 5; Conservative 5; Mismatches 1; Indels

;

707 kfnrcikrkve 717 원

| ::|:::||| 8 KSSKCVRQKVE 18

ð

Search completed: Wed Jan 28 12:09:58 1998 Job time : 14 secs.

Jan 28 12:22

| (TM) | ****** |
|----------------|---|
| | ****** |
| | ***** |
| | ****** |
| | ********* |
| | ********* |
| | ********* |
| | ******* |
| <u>E</u> ===== | *************************************** |

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 University of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

protein - protein database search, using Smith-Waterman algorithm MPerch_pp

Wed Jan 28 12:14:09 1998; MasPar time 57.52 Seconds 887.795 Million cell updates/sec Run on:

Tabular output not generated.

Title:

>US-08-487-283A-2 (1-1676) from US08487283A.pep 12048 Description: Perfect Score:

1 MGLLGILCFLIFLGKTWGQE.......CQAFLANLDEFAEDIFLNGC 1676 Sequence:

Scoring table:

PAM 150 Gap 11

95051 seqs, 30469580 residues Searched:

Post-processing: Minimum Match 0% Listing first 45 summaries

Database:

1:ann1 2:ann2 3:ann3 4:ann4 5:unann1 6:unann2 7:unann3 8:unann4 9:unann5 10:unann6 11:unann7 12:unann8 13:unann9 14:unann10 15:unann11 16:unann12 17:unenc

Mean 56.253; Variance 125.273; scale 0.449

Statistics:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| Pred. No. | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | _ | _ | _ | _ | 0.00e+00 | 0.00e+00 | 0.00e+00 |
|--------------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|---|
| Description | complement C5 precur | complement C5 precur | complement C3 precur | complement C3 precur | complement component | complement C3 precur | complement C3 precur | complement C3 precur | complement C3 precur | cobra venom factor p | complement protein h |
| iD | СБНО | C5MS | C3NJ | СЗНО | 150806 | 150711 | C3MS | C3GP | C3RT | 151018 | 521045 |
| 80 | 7 | 7 | 7 | 7 | 15 | 9 | ~ | ~ | ~ | ڡ | 15 |
| Query Match Length DB | 1676 | 1680 | 1651 | 1663 | 1673 | 1652 | 1663 | 1666 | 1663 | 1642 | 1620 |
| Query Match | 100.0 | 81.3 | 19.6 | 19.6 | 19.2 | 19.1 | 19.1 | 19.1 | 18.7 | 18.4 | 17.5 |
| Score | 12048 | 9789 | 2367 | 2361 | 2309 | 2298 | 2296 | 2302 | 2255 | 2214 | 2106 |
| Result No. | 1 | 7 | ٣ | 4 | S | 9 | 7 | œ | 6 | 9 | ======================================= |

| **** |
|----------------------|
| ***** |
| **** |
| ***** |
| **** |
| ***** |
| ***** |
| **** |
| |
| **** |
| **** |
| |
| ಿ |
| |
| |
| |
| |
| - |
| - |
| |
| • |
| C |
| |
| |
| : 800 |
| |
| |
| • |
| - |
| |
| |
| |
| |
| |
| - |
| US-08-487-283A-2.rpt |
| _ |
| _ |
| _ |
| _ |
| 7 |
| _ |
| - |
| ~ |
| ₩ |
| 7 |
| <u> </u> |
| _ |
| |
| 7 |
| 7 |
| 7 |
| 7 |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| Jan 28 12:22 (|

| | | human sion 30-Sep-1992 ftext_change 11266; S15121 J.C.; Fleischer, D.T.; Hunt, A.; -368 human complement pro-C5. Evidence derived from a single copy gene. d d 4-1482 n complement component C5: ctural gene to chromosome 9. |
|--|--|--|
| 8 | 42210 33329 20872 20872 41185 41185 55144 4102 48771 6188 6188 6188 6188 6188 6188 6188 | rte has the hard the |
| 1620 1744 1735 1735 1735 1736 1736 1487 1477 1477 1477 1476 | 1500 | U plement anaphyl srmal nam srmal nam srmal nam 3-192 075, A27 075, A27 075 iland, D iland, D |
| | 25 525 26 506 27 507 28 493 30 470 31 456 33 349 34 370 35 349 36 354 37 319 38 267 40 255 41 251 42 235 44 235 | RESULT 1 C5H TITLE COM CONTAINS C5a CONTAINS #fo DATE 30- ACCESSIONS A40 REFERENCE A40 # journal J, W # journal J, W # journal J, W # fittle Com # cross-reference # accession A40 # fross-reference # accession A40 # fross-reference # accession A40 # fournal Biologule_typ # fross-reference # accession A40 # fournal Biologule_typ |

Inn 28 12-22

#authors

REFERENCE

journal

title

m

contents

COMMENT

678-1676

19-673

GENETICS

COMMENT

KEYWORDS

*accession

#authors

REFERENCE

journal

title

##note

fauthors

REFERENCE

iournal

#title

US-08-487-283A-2.pr Jan 28 12:22

#product C5b alpha' chain #status predicted #label C5BA\ ö product C5a anaphylatoxin #status experimental #label 61 dkkfsyssghvhissenkfgnsailtigpkglpgggppvsyvylevvskhfskskrmpit 120 121 ydngflfihtdkpvytpdqsvkvrvyslnddlkpakretvltfidpegsevdmveeidhi 180 181 giisfpdfkipsnprygmwtlkakykedfsttgtayfevkeyvlphfsvsiepeynfigy 240 361 LFLKPGIPYPIKVQVKDSLDQLVGGVPVILNAQTIDVNQETSDLDPSKSVTRVDGVASF 420 61 DKKFSYSSGHVHLSSENKFQNSAILTIQPKQLPGGQNPVSYVYLEVVSKHFSKSKRMPIT 120 300 300 361 lflkpgipypikvqvkdsldqlvggvpvilnaqtidvnqetsdldpsksvtrvddgvasf 420 421 vlnlpsgvtvlefnvktdapdlpeengaregyraiaysslsgsylyidwtdnhkallvge 480 481 hlniivtpkspyidkithynyliiskgkiihfgtrekfsdasyqsinipvtqnmvpssrl 540 HINIIVTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPSSRL 540 541 lvyyivtgeqtaelvadsvwlnieekcgnqlqvhlapdadayspgqtvslnmatgmdawv 600 GIISFPDFKIPSNPRYGMWTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFIGY 240 fbinding_site carbohydrate (Asn) (covalent) fstatus #binding_site carbohydrate (Asn) (covalent) #status Gaps #length 1676 #molecular-weight 188330 #checksum 3858 #cleavage site Arg-Leu (C5 convertase) #status experimental\ ö Length 1676; 0; Indels disulfide_bonds #status predicted/ 100.0%; Score 12048; DB 2;

| | ••• |
|----------------------------|-----|
| | 98 |
| | |
| | |
| ***** | • |
| | |
| ***** | ж |
| 355565 | ٠. |
| | ••• |
| | |
| | • |
| | |
| | ••• |
| | |
| | |
| | • |
| | |
| | |
| | 33 |
| 00000 | ••• |
| | 99 |
| ***** | ** |
| | ** |
| 20000 | • |
| | |
| | ж |
| 99999 | 33 |
| | |
| | |
| | |
| | |
| | |
| | |
| ••••• | |
| 355665 | 88 |
| | • |
| | |
| ***** | • |
| | 88 |
| | ** |
| | |
| | ٠. |
| | |
| | ••• |
| | 99 |
| | |
| | |
| | • |
| | |
| | ж |
| 273 | • |
| | |
| | |
| ∞ | 88 |
| 56 | × |
| × | |
| × | |
| ౙ | |
| 83 | |
| 83 | |
| 83 88 | |
| 82-58 | |
| 487-28 | |
| 487-28 | |
| 487-28 | |
| 8487-28 | |
| 8487-28 | |
| 88.78 | |
| -08-487-28 | |
| -08-487-28 | |
| S-08-487-28 | |
| IS-08-487-28 | |
| JS-08-487-28 | |
| US-08-487-28 | |
| US-08-487-28 | |
| US-08-487-28 | |
| US-08-487-283A-2.rpr | |
| US-08-487-28 | |
| 87-68-487-28 | |
| US-08-487-28 | |
| 87-187-83 112-08-487-28 | |
| US-08-487-28 | |
| | |
| | |
| | |
| | |
| | |
| 2:22 US-08-487-28 | |

S

| | | | 8 (|
|----------|------|--|------|
| ð | 541 | . LVYYIVTGEQTAELVSDSVWI.NIEEKCGNQI.QVHI.SPDADAYSPGQTVSI.NMATGMDSWV | 009 |
| QG | 601 | alaavdaavygvqrgakkplervfqfleksdlgcgaggglnnanvfhlagltfltnanad | 099 |
| ò | 601 | ALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGINNANVFHLAGLTFLTNANAD | 099 |
| g | 661 | dsqendepckeilrprrtlqkkieeiaakykhsvvkkccydgacvnndetcegraaris | 720 |
| ð | 661 | - 8 | 720 |
| g | 721 | gprcikafteccvvasqlranishkdmqlqrlhmktllpvskpeirsyfpeswlwevhlv | 780 |
| ò | 721 | | 780 |
| g | 781 | | 840 |
| à | 781 | PRRKQLQFALPDSLTTWEIQCIGISNTGICVADTVKAKVFKDVFLEMNIPYSVVRGEQIQ | 840 |
| q | 841 | lkgtvynyrtsgmgfcvkmsavegictsespvidhggtksskcvrgkvegssshlvtf | 006 |
| ð | 841 | | 006 |
| g | 901 | Ipleiglhninfeletwfgkeilvktlrvvpegvkresysgvtldprgiygtisrrkefp | 096 |
| à | 901 | LP LEIGLHININFSLETWFGKEILVKTLAVVPEGVKRESYSGVTLDPRGIYGTISRRKEFP | 096 |
| QQ | 961 | yripldlypkteikrilsvkgllygeilsavlsqeginilthlpkgsaeaelmsvypvfy | 1020 |
| õ | 961 | YRIPLDLVPKTEIKRILSVKGLLVGEILSAVLSQEGINILTHLPKGSAEAELMSVVPV | 1020 |
| qo . | 1021 | vfhyletgnhwnifhsdpliekqklkkklkegmlsimsyrnadysysvwkggsastwlta | 80 |
| ð | 1021 | VFHYLETGNHMNIFHSDPLIEKQKIKKKIKEGMLSIMSYRNADYSYSVWKGGSASTWLTA | 1080 |
| g | 1081 | falrulgqvnkyveqnqnsicnslwlvenyqldngsfkensqyqpiklqgtlpvearen | 1140 |
| δ | 1081 | FALRVI.GQVNKYVEQNQNSI.CNSI.LMLVENYQL.DNGSFKENSQYQPIKI.QGTI.PVEAREN | 1140 |
| 요 | 1141 | slyltaftvigirkafdicplvkidtalikadnfllentlpagstftlaisayalslgdk | 1200 |
| ٥y | 1141 | | 1200 |
| qq | 1201 | thpqfrsivsalkrealvkgnppjyrfwkdnlqhkdssvpntgtarmvettayalltsln | 1260 |
| à | 1201 | | 1260 |
| DP DP | 1261 | lkdinyvnpvikwlseeqrygggfystqdtinaieglteysllvkqlrlsmdidvsykhk | 1320 |
| ογ | 1261 | IKDINYVNPVIKWLSEEQRYGGGFYSTQDTINAIEGLT3YSLLVKQLRLSMDIDVSYKHK | 1320 |
| g | 1321 | galhnykmtdknflgrpvevllnddlivstgfgsglatvhvttvvhktstseevcsfylk | 1380 |
| δ | 1321 | GALHNYKMTDKNFLGRPVEVLIADDLIVSTGFGSGLATVHVTTVVHKTSTSEEVCSFYLK | 1380 |
| e e | 1381 | idtqdieashyrgygnsdykrivacasykpsreesssgsshavmdislptgisaneedlk | 1440 |
| ολ | 1381 | IDTQDIEASHYRGYGNSDYKRIVACASYKPSREESSSGSSHAVMDISLPTGISANEEDLK | 1440 |
| DP DP | 1441 | alvegvdqlftdyqikdghvilqlnsipssdflcvrfrifelfevgflspatftvyeyhr | 1500 |
| ý | 1441 | ALVEGVDQLFTDYQIKDGHVILQLASIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHR | 1500 |
| Ор | 1501 | pdkqctmfystsnikiqkvcegaackcveadcgmqeeldlisaetrkqtackpeiaya | 1560 |

Jan 28 12:22 US-48-487-283A-2.7pt

1501 PDKQCTMFYSTSNIKIQKVCEGAACKCVEADCGQMQEELDLTISAETRKQTACKPEIAYA 1560

ð

Ø

Haviland, D.L.; Haviland, J.C.; Fleischer, D.T.; Wetsel, R.A. J. Biol. Chem. (1991) 266:11818-11825 Structure of the murine fifth complement component (C5) gene. A large, highly interrupted gene with a variant donor splice site and organizational homology with the third and fourth complement component genes. 22/3; 86/3; 140/3; 164/3; 195/2; 223/1; 253/2; 291/3; 334/1; 372/3; 434/3; 502/3; 572/3; 622/3; 667/1; 691/1; 757/1; Wetsel, R.A.; Fleischer, D.T.; Haviland, D.L. J. Biol. Chem. (1990) 265:2435-2440 Deficiency of the muxine fifth complement component (C5). A anaphylatoxin from the amino end of the alpha chain, generating Activation of C5 initiates the spontaneous assembly of the late complement components, C5-C9, into the membrane attack complex. C5b has a transient binding site for C6. The C5b-C6 complex is complement C5 precursor - mouse C5a anaphylatoxin, C5b #formal name Mus musculus #common_name house mouse 19-Nov-1988 #sequence_revision 15-Oct-1994 #text_change Complement C5 contains two disulfide-linked chains, formed by removal of four basic residues. C5 convertase releases C5a the foundation upon which the membrane attack complex is Wetsel, R.A.; Ogata, R.T.; Tack, B.F. Biochemistry (1987) 26:737-743 Primary structure of the fifth component of murine C5a has potent spasmogenic and chemotactic activity. 2-base pair gene deletion in a 5'-exon tross-references MUID:90153853 ##residues 1-215,'L' ##label WET ##cross-references GB:J05234 #type complete C5b (beta and alpha' chains). A35530; A27538; A40429 1-15 ##label HAV cross-references MUID:87185363 fcross-references MUID:91268053 f#cross-references GB:M64852 16-Feb-1997 .complement ##molecule_type mRNA ##molecule_type DNA A27538 A40429 assembled C5MS #map_position 2 ##residues #accession #accession accession REFERENCE #authors authors journal journal #authors | journal **ACCESSIONS** REFERENCE #title title REFERENCE title CONTAINS ORGANISM GENETICS COMMENT COMMENT RESULT ENTRY COMMENT TITLE 음 g ð 8

CLASSIFICATION

KEYWORDS FEATURE 679-1679

156-1679 679-755

481 541 LLL 541 1137 099 837 961 1197 1381 601 601 721 781 841 901 1077 1141 1201 1261 1257 1317 661 1081 염 염 പ്പ 유 g ò ð 염 9 g g g 염 g g a g ð à ò ò ò δ Ş ð δ à ð à ð à g glycoprotein; inflammation; membrane attack complex; plasma C5a anaphylatoxin #status predicted #label C5TN C5b alpha' chain #status predicted #label C5BAN ; 787/2; 812/1; 858/3; 934/3; 955/1; 985/1; 1056/1; 1081/2; 1134/3; 1166/3; 1224/1; 1292/3; 1343/3; 1364/3; 1392/1; 1411/2; 1445/3; 1410/3; 1506/1; 1534/1; 1564/1; 1592/1; dkkvtfssgyvnlspenkfqnaalltlqpnqvpreespvshvylevvskhfskskkipit 120 120 240 240 300 300 360 360 361 lfvkpgipfsikaqvkdslegavggvpvtlmaqtvdvnqetsdletkrsithdtdgvavf 420 420 480 #binding site carbohydrate (Asn) (covalent) #status predicted 8 9 Gaps mglwgilcllifldktwggeqtyvisapkilrvgssenvviqvhgyteafdatlslksyp knfknfeitvkaryfynkvvpdaevyaffglredikdeekqmmhkatqaaklvdgvaqis fdsetavkelsynsledlnnkylyiavtvtessggfseeaeipgvkyvlspytlnlvatp giisfpdfkipsnpkygvwtikanykkdftttgtayfeikeyvlprfsvsielertfigy 421 vlnlpsnvtvlkfeirtddpelpeenqaskeyeavaysslsqsyiyiawtenykpmlvge 7; Mismatches 149; Indels Length disulfide_bonds #status predicted\ #length 1680 #molecular-weight 188876 9789; DB 2; No. 0.00e+00; Score Pred. 218; n 81.3%; Similarity 77.8%; tches 1307; Conservative 1637/2 19-674, 679-1679 19-674, 756-1679 19-674 702-728, 703-735, 715-736, 870-1531, 567-814, 635-670 915, 1119, 1633

Query Match Local

SUMMARY

61 19 171 121

유 Š g ð 셤 241 241 301

용 ð 셤 3 셤

301

361

õ

ď

181

ð

181

g

ð

1105-1163, 1379-1509, 1409-1478, 1524-1529, 1536-1609, 1557-1679,

US-08-487-283A-2.rpi

1020 1016 1080 1076 1140 1136 1260 1256 1320 1316 1200 fylkidtqdieasshfr-lsdsgfkriiacasykpskeestsgsshavmdis1ptgigan 1439 480 900 900 99 629 780 776 840 836 900 968 960 926 421 VINIPSGYTVILEFNVKTDAPDI.PEENQAREGYRAIAYSSI.SQSYLYIDWTDNHKAII.VGE 481 ylnimvtpkspyidkithynylilskgkivqygtreklfsstyqninipvtqnmvpsarl lvyyivtgeqtaelvadavwinieekcgnqlqvhlspdeyvyspgqtvsldmvteadswv LVYYIVTGEQTAELVSDSVWINIEEKCGNQLQVHLSPDADAYSPGQTVSLNMATGMDSWV alsavdravykvqgnakramqrvfqaldeksdlqcqaqqqhdnadvfhlagltfltnana rvtigplcirafnecctiankirkesphkpvqlgrihiktllpvmkadirsyfpeswlwe ihrvpkrkqlqvtlpdslttweiqqigisdngicvadtlkakvfkevflemnipysvvrg eqiqlkgtvynymtsgtkfcvkmsavegictsgssaaslhtsrpsrcvfqriegssshlv tftllpleiglhsinfsletsfgkdilvktlrvvpegvkresyagvildpkgirgivmrr kefpyripldlvpktkverilsvkgllvgeflstvlskeginilthlpkgsaeaelmsia aqektlyltafsvigirkavdicptmkihtaldkadsfllentlpskstftlaivayals lgdrthprfrlivsalrkeafvkgdppiyrywrdtlkrpdssvpssgtagmvettayall aslklkdmnyanpiikwlseeqrygggfystqdtinaieglteyslllkqihldmdinva ykhegdfhkykvtekhflgrpvevelnddlvvstgyssglatvyvktvvhkisvseefcs

8 a õ a

1500

1560

g

ð

1619

셤 õ 셤 ð

Š

O,

A46513 A46513

ACCESSIONS

ORGANISM CONTAINS

ENTRY TITLE REFERENCE

authors f journal

title

A46513

f residues accession

fnote

##note

COMMENT

COMMENT COMMENT

US-08-487-283A-2.rpr Jan 28 12:22

predicted #label C3T\ pathway; glycoprotein; hydrolase; immune response; inflammation; liver; plasma; serine proteinase; thiolester bond 97; fproduct complement C3b alpha' chain #status predicted fproduct complement C3 #status predicted #label CC3\
fproduct complement C3b #status predicted #label C3B\ major site of synthesis of this plasma protein is the liver. domain signal sequence fstatus predicted flabel SIGN fproduct complement C3 alpha chain #status predicted 121 lsyqsgfvfiqtdkgiytpgspvryrvfsvdhnmhrmdktvivefqtpegivvs-skpvn 179 prkqktlfqsrvdmnqagsmfvtpti-kvpakelnkdskqnqyvvvkvtgpqvalekvvl 120 psgsir-p-ynlpelvsfgtwkavakyehspeesytayfdvreyvlpsfevrlqpsdkfl 237 179 HIGIISFPDFKIPSNPRYGM#TIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFI 238 eatlkrdt-lrs-rfqdlnqlvghtlyvsvtvitesgsdmvvteqggihivtspyqiyft 350 238 -yidgnknfhvsitarylygkkv-egvafvvfgvk--i-ddakksipdsltripiidgdg 292 358 ATPLELKPGIPYPIKVQVKDSLDQLVGGVPVILNAQTIDVNQETSDLDPSKSVTRVDDGV 417 19 Gaps cross-link thiolester (Cys-Gln) #status predicted product complement C3 and C3b beta chain fstatus 4 malylvaalligfpg-s-shgalytlitpavlrtdteeqilveahgdstpksldifvhdf 60 PDKKESYSSGHVHLS-SENKEQNSA1LTIQPKQLPGGQNPVSYVXLEVVSKHFSKSKRMP QVTFDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLALV 351 ktpkyfkpgmpyeltvyvtnp-d---gs-paa-hvp-v-vs-ea--ih-segtt-lsdgt #cleavage \bar{s} ite Arg-Ser (C3 convertase) #status predicted\ #length 1651 #molecular-weight 184925 #checksum 518; Conservative 416; Mismatches 655; Indels 127; DB 2; Length 1651; fsuperfamily alpha-2-macroglobulin acute phase; complement alternate pathway; disulfide_bonds fstatus predicted/ C3a anaphylatoxin #status Query Match 19.6%; Score 2367; DB 2; Best Local Similarity 30.2%; Pred. No. 0.00e+00; fregion properdin binding\ predicted #label C3BB\ ð

Jan 28 12:22

셤

g

õ 염 õ g õ a õ

Š

ΩP

Š Ωp õ g

 \equiv

Jan 28 12:22

S G

S d

9 9

õ 셤 g

õ

Ð

\$

ద

õ

셤

8

õ

US-08-487-283A-2.rpt

| Jan 28 12:22 | US-08-487-283A-2-pr |
|--|---|
| #journal #title | J. Biol. Chem. (1975) 256:8293-8301 Human anaphylatoxin (C3a) from the third component of |
| #cross-refere | |
| ##CCESSION ##molecule ##regidnes | |
| REFERENCE | A27603 |
| #authors | Daoudaki, M.E.; Becherer, J.D.; Lambris, J.D. J. Tamings, (1988) 140-1577-1580 |
| # journar #title | A 34-amino acid peptide of the third component of complement |
| forose_refere | mediates properdin binding. |
| #accession | A27603 |
| ##molecule ##residues | ##molecule type protein |
| REFERENCE | A23435 |
| fauthors | Hellman, U.; Eggertsen, G.; Engstrom, A.; Sjoquist, J. Biochem, T. (1908), 930, 252, 261 |
| t journar ftitle | Proceedings of (1907) 2333-331 Amino acid sequence of the trypsin-generated C3d fragment |
| | from human complement factor C3. |
| #Cross-relered | nces Mulliabuza442 A23435 |
| ##molecule | type protein |
| ##residues | 1002-1012, E', 1014-1303 ##label HEL |
| ##note | sequence corresponding to residues 10/2-1100 was not determined but was taken from de Bruijn & Fey |
| PEFFENCE | (reference A946U5) |
| fauthors | Poznansky, M.C.; Clissold, P.M.; Lachmann, P.J. |
| # journal | |
| #title | between human C3F and C3S results |
| | single amino acid change from an asparagine to an aspartate residue at position 1216 on the alpha-chain of the |
| | lement component, C3. |
| #accession | A45830 |
| ##Bratus ##molecule | not compared with conceptual translation type DNA |
| ##residues | 1212-1215, N', 1217-1223 ##label P02 |
| ##note | tnis is the USS allele R45830 |
| ##status | not compared with conceptual translation |
| ##molecule | type DNA |
| ##residues | 1212-1223 ##label PO2 |
| her enember | Dolmer. K.: Sottrup-Jensen. L. |
| # journal | 315:85-90 |
| forter | Distillate bilages in numan complement component com. empotation: disulfide bonds |
| þ | sequence shown is the C3 fast (C3F) allele, which is found |
| ien fai | |
| COMMENT COMPL | C3 contains two cha |
| | J. L J. 44 |

C3b, with its highly reactive thiol group, binds to the surface of foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the

residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and alternative complement pathways, releases the C3a anaphylatoxin

from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form the alternative-complement-pathway C3/C5 convertase.

C3a anaphylatoxin is a vasoactive peptide and a mediator of inflammation.

COMMENT

셤

유

ŝ

Š

임

g

Š

δ

음

õ

g

Š

셤

셤

ð

2

| <pre>Db 1050 kkgytqqlafrqpssafaafvkrapstwltayvvkvfølavnliaidsqvlcgavkwlil 1109</pre> | Db 1110 ekqkpdgvfqedapvihqemigglr-nnnekdmaltafvlislqeakdiceeqvnslpgs 1168 : : : | Db 1169 itkagdfleanymnlqrsytvaiagyalaqmgrlkgpllnk-flttakdk-nr- 1219 : : : : : | Db 1220 w-ed-pgkg-l-ynveatsyallallqlkdfdfvppvvrwlnegryygggyst 1269 :: :: :: | Db 1270 qatfmvfqalaqyqkdapdhqelnldvalqlpsraskithrihwesasllrseetkeneg 1329 | Db 1330 ftvtae-gkgggtlsvvtmyhakakdgltcnkfdlkvtikpapetekrpgdakntmilei 1388 : : : : : : | Db 1389 ctryr-g-dqdat-msildismmtgfapdtddlkqlangvdryiskyeldkafsdrnt 1443 : : :: : :: : ::: : : : :: :: Qy 1405 CASYKPSREESSSGSSHAVMDISLPTGISANEEDLKALVEGVDQLFTDYQI-KD-GH- 1459 | Db 1444 liiyldkvshseddclafkvhqyfnveliqpgavkvyayynleesctrfyhpekedgkln 1503 : : : : : : : : : | Db 1504 klcrdelcrcaeencf-iqksdd-kvtleerldkacepgvdyvyktrlvkvqlsndfdey 1561 : : : | Db 1562 imaieqtiksgsdevqygqrtfispikcrealkleekkhylmwglss-df-wgekpnls 1619 :: | Db 1620 yiigkdtwvehwpeedecqdeenqkqcqdlgaftesmvvfgc 1661 | RESULT 5 150806 #type fragment | | rs al | eren | #accession loudub ##status preliminary; translated from GB/EMBL/DDBJ ##molecule_type_mRNA ##residins 1-1673 ##label NON |
|---|---|---|--|---|--|---|---|--|---|--|--|---|---|--|---|
| 124 slqsgylfiqtdktiytpgstvlyriftvnhkllpvgrtvmvnienpegipvkqdslssq 183 : ::! : : : ::: :: : : 120 TYDNGFLFIHTDKPVYTPDQSVKNRVYSLNDDLKPAKRETVLFIDPEGSEVDMYEEI 177 | 184 nqlqvlpls-wdipelvnmgqwkirayyenspqqvfstefevkeyvlpsfeviveptekf 242 ::: ::::: | 243 yyi-yne-kglevtitarflygkkv-egtafvifgi-qdgegrislpeslkripiedg 296 | 297 sgevvlsrkvildgvantraedlvgkelyvsatvilhsgsdmvqaersgipivtspyqih 356 :: : :: :: 296 IAQVTFDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLN 355 | 357 ftktpkyfkpgmpfdlmvfvtnp-dgs-pay-rvp-vav-qge-dtvqsltq-gd 404 | 405 gvaklsinthpsqkplsitvrtkkqelseaeqatrtmqalpystvgnsnnylhlsvlrte 464 : :: : : : : : : | 465 lrpgetlnvnfllrmdraheakiryytylimnkgrllkagrqvrepgqdlvvlplsittd 524 : :: : : : :: : : : : : | 525 fipsfrlvayytligasggrevvadsvævdvkdscvgslvvksggsedrgpvpggqmtlk 584 | 585 iegdhgarvvlvavdkgvfvlnkknkltqskiwdvvekadigctpgsgkdyagvfsdagl 644 : :: : : : :: : : : | 645 tftsssggqtagraelgcpgpaarrrrsvgltekrmdkvg-kypkel-rkccedgmrenp 702 | 703 mríscgrittíislgeackkvíldccnyitelriqharashlglarsnldedijaeeniv 762 : :: : : :: :: ::: :: | 163 srsefpeswlwnvedlkeppkngistklmniflkdsittweilavsmsdkkgicvadpfe 822 :- :- 1111 765 IRSYFPESWLWEVH-L-VPRRKQL-QFALPDSLTTWEIQGIGISNT-GICVADTVK 816 | 823 vtvmqdffidlrlpysvvrneqveiravlynyrqnqelkvrvellhnpafcslattkrrh 882 :::: ::::: : 817 AKVFKDVFLEMNIPYSVYRGEQIQIKGTVYNYRTS-GHQFCVKMSAVECICTSESPVIDH 875 | 883 qqt-vtippk <i>ssls</i> vpyvivplktglqevevkaavyhhfisdgvrkslkvvpeg 935 : : : :: ::: 876 QCTKSSKCVRQKVEGSSSHLVTFTVLPLEIGIHNINFSLETWFGKEILVKTLRVVPEG 933 | 936 irmnktvävrtldperl-gregvqkedip-padlsdqvpctesetril-lqgtpvagmte 992 :: : : : | 993 davdaerlkhlivtpsgcgeqmmigmtptviavhyldeteqwekf-qlekrggaleli 1049 : :: : |

õ 음 Š 원 Š 셤 õ ä ò 유 õ В 8 셤 õ

유

Š

SUMMARY

1 vl-llmsvgtsvtqdpmvllsvpsviligsdvnvlvdhaastedvsvvvraeefltkk-q 58 Gaps 458; Mismatches 636; Indels 130; Length 1673; Score 2309; DB 15; Pred. No. 0.00e+00; 19.2%; 29.5%; 512; Conservative Local Similarity Query Match Matches Best 용

> ð 9

8

latqtitltqldpaiatlklgfdienpdktnsastkhhvrlv-akvesksfnkeitahal
:: : : : : : : | | | | | | :
YSSGHVHLSS-ENKFQNSAI-LTIQ-P-KQLPGG-QNPVSYVYLEVVSKHFSKS-K-RMP 99

lsyrsghvvvqtdkpiytpdekvkyrmfpmnredvhripvrqsmtvdivnadgviverqi 177 118 9 ò

237 ktikatdegivdgtsftipaiskhgtwkifarmsgapninssaefdvreyilptfevkin 178 g 8

EEI-D-HIGIISFPDFKIPSNPRYGMFTKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIE 232 175 290 pkqrvfhind-eefvvditanyfnqelvsgta-yvryfl-en-gdvpk-lvdss--sttl 238

9 Š 셤 Š 9 Š

233

291

352 293

qitatktsryfkpelpyfiqvevrnadgspskevdvv--ak-vqvgsat--inpqk-m-r 401 349

: |:::|| :|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:||

tdsngltsftvt-ppnvngltvtvrtderh-psneggelvytagkyas-a-symhid-v- 455 VD-DGVASFVIALIPSGVTVLEFNVKTDAPDLPEENQAREGYRAIAYSSLSQSYLYIDWTD 471 :: 413 402 g ð

514 t-rimrlgetlnvfltakttqlnavthftymvltrqvivktnrktkesgggpsnvripit
:::|| ||:::||:|:::||:||:||:||:|||:|||||
NHKALLVGEHINIIVTPKSPYIDKIŢHYNYLILSKGKIIHFGTREKFSDASYQSINIPVT 456 g

| pdmaprfrflayyilpg---geivadsvtvevtelcksqvslslkgrp-tlepkamltld 570 472 515 В õ

630 591 571 ligepdarvgllavdgavyavnrkhrltgdrvwkametfdtgctaeggagrpgvfsdagl QNMVPSSRLLVYY I VTGEQTAELVSDSVWLN I EEKCGNQLQVHLSPDADAYSPGQTVSLN 532 g ò

592

8

9

õ

651

qgceerlkrvt-gpkecvdaflqcckkaeeyrkseslgaktvlrrndfmeldlmnedevn 069 709 g ð

809 tkdffiklhlpysvkrgeqteirvilynymeesltiltemdivesicst-sk--s--gak

g

US-08-487-283A-2.rpi

∞

1205 1262 1320 1379 1439 1448 1499 1292 1622 1031 1091 1401 dnrtevgsfvyytvniqtviksgqdqaiqpkatrl-fivtrscdgrlgmetprqyllmgr 1617 879 974 995 kg-etk-drndrfqyvldasswveqwpvdekcnqpnvqtfcaikreyefsmqiqgc 1671 : : | : | | | | : : | | | | : : | | EALQIKYNFSFRYIYPLDSLTWIEYWPRDTTCS--SCQAFLANLDEFAEDFFLNGC 1676 di-r--s-esrsvh-veeretffikneispdvvpnsdvltfisvkgdelaetmvncldak : ::|:: | | |: |:||| ::||| ::| : | | ESYSGVTLDPRGIYGTISRRKEF-PYRIPLDLUPKTEIKRILSVKGLLVGEILSAVLSQE elsyrkadhsyaafikrpsstwltafvvkvyslakrviivdnqelcgpvewiiknrqnsd gsyredgpvihremqgg--vggteghvsmtafiligiqqaqeycgvsvpnykqsmnravq algamagykmdssskel-idvgleit-spknnfekkmkiteetrfvgephkippggniti yisnyeitdsvliiywdkvpstedycfafkikqmlrsdmiqpvtasvydyyspadkctrl ynlpggyvelsplcqndlcqcvevscpakkpkfdtsitvlhrqeaacvagidyay-vgiv | || || : : || : || : || : || :| || SITVE-NVFVKYRATLLDIYKTG-E-AVAEKDSEITFIKKVTCTNA-ELVKGRQYLIMGK flaskvsdlkrmytiaitryalalqd---peseaahsswkklenrttfeskghrywkae-1116 1174 1293 1321 1618 1263 1349 1449 1500 1509 1559 864 880 920 937 966 1150 1402 1440 1623 975 1032 1092 1380 1567 5 셤 ð 유 ð 유 ð 셤 à 연 ð 임 8 9 5 임 8 g ð g ð 유 ð 셤 셤 ð a ð ð

chicken formal_name Gallus gallus #common_name precursor - chicken #type complete complement C3 TITLE ORGANISM RESULT ENTRY

Jan 28 12:22

Jan 28 12:22

US-08-487-283A-2.mr

| 590 IAMATGMDSWVALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGLANANVFHLA 649 | 635 glsltsnvmintegrsevgcakpakrkrrsvrlikhkgtkmaeysdknlrkccedgirkn 694 | DSQENDEPC-KEIL | 695 lmgyscekratyv-ldakscteaflscclyikgirdeerelgyelarsevddaflsdedi 753 : : : : : : 1052CVVASQLRANISHKDMQLGRLHMK-TLLPVSKP 763 | 754 tsrslfpeswiwqveelteppneggismkljpjylkdsittwevlavsisenkglcvadp 813 24 | | 871 -k-tryqq-ifq-lepqssdavpfvivplelgqhdvevkaavwnsfvsdgvkklrvvpe 926 : : | 927 gmrlektvkiveldpktl-gnngvq-evkvkaanlsdivpntesetkvsiggnpvs-ilv 983 :: : | 984 ekatdgtklkhlivtpsgcgeqmnigmtptviavhyldstmqwetfginrrteaielikk 1043 :: :: | 1044gytqqlayrkedgsfaafttrpsstwltayvakvfamainmvdikpevvcgaikwli 1100 :: : : :: :: :: :: :: : : | 1101 lekqqpdglfqedapvihkemvggyh-ga-epsvsltafvlsalqesqkicknyvksldg 1158 | 1159 siakasdylerkyqsltrpytvaltsyalaltgklnse-kvlmkfsk-dgth | 1209 -wa-e-rnaht-yniegtsyalvallqmekaeltgpvvrwlaqqnyfgggygs 1257 ::::: : : : : : : : : : ::::: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: :: ::: :: ::: ::: ::: ::: ::: ::: ::: ::: ::: :: ::: ::: ::: :::: | 1258 tqatilvfqalaqyhvalprhvelnldvsvllprranaityriennnalvarsaetklne 1317 :: : : :: :: :: | 1318 dftvkae-gtgkgtmtvvtyykakvpekenkcdnfdlrvsvedvkagreve-gvirsvki 1375 : : : : | 1376 tictrf-ldtvdatm-sildismltafspdvqdlkslsegveryiskfeidhalenr 1430 : : :: :: :: : : : | 1431 snliiyldkvshqveeciafrahqhfqvqliqpasvivysyykiddrctrfyhpdkaggq | 1491 Irkichgevc-caeencf-irvkkdnpitvneridlackpgvdyvykvkvvateetpshd 1548 |
|--|--|----------------|--|---|---|--|---|---|---|--|---|---|---|--|--|---|--|
| ð | g G | ò | d y | g : | \$ 6 \$ | g & | අ දු | e & | 음 & | g Vy | අ දි | g & | Up Qy | qa Vy | DP Qy | Oy Oy | Op |
| DATE 13-Sep-1996 #sequence_revision 13-Sep-1996 #text_change | ACCESSIONS 150711 | rB | fittle Isolation, primary structure, and evolution of the third component of chicken complement and evidence for a new member of the alpha 2-macroglobulin family. | <pre>#cross-references MUID:95173423 #accession I50711 ###tataur ###tataur ##statur ##statur</pre> | ##factoruse_type mknn ##residues 1-1552 ##label MAV ##cross-references EMBL:016848; NID:g755814; CDS_PID:g755815 CLASSIFICATION #superfamily alpha-2-macroglobulin SUMMARY #length 1652 #molecular-weight 184085 #checksum 6420 | Query Match 19.1%; Score 2298; DB 6; Length 1652; Best Local Similarity 29.1%; Pred. No. 0.00e+00; Matches 502; Conservative 455; Mismatches 645; Indels 122; Gaps 91; | Db 1 mgll-llplllgvlllhavptpaqmvtmvtpavlrldtdekvvleapglsapteanilvq 59 | Db 60 dfpqkrkvlfqvrkqlnpaegmmaiatvkvpvkllpp~~ygk-hfvsvvarvgqvtlekvl 118 : : : :: : | Db 119 lvslqsqhif1qtdkpiytpgstvlsrlfalshfmqpllktvivevktpdnviikqvpvs 178 :::: ::: : : ::: ::: ::: ::: ::: ::: ::: ::: ::: ::: | <pre>Db 179 spmrngifsi-nhnlpevvslgtwtitakfedsqdqvfstqfevkeyvlpsfevtldpqe 237 : : : : :: </pre> | Db 238 kfl-yidpaedfrvtitarylygknl-ggtafvlfgvvvddekktipgs-lgrvkvtd 292 | <pre>Db 293 gdgqavl-pmamlrq-pfanlqelvghalyvtvtvltesgsdmveaqrsgirivtspyti 350 i i : i : i : i : i </pre> | Db 351 hfthtpkyfkpgmpfdptvyvtnp-dns-paaa-gipvkadnfqglvstqr 398 :: :: : :: | Db 399 dgtaklvinmpanknsvpitvrtdqkdlpperqaerqivaeayqsqgnsgnylhlavgas 458 | Db· 459 qvqpqnlpinfhlksnrddvrksvsyftylilskghivhvgrqpregdqslvtmslpvt 518 : : : : : : | <pre>Db 519 anlipsfrivayyhvkpgeiiadsvwvdvkdtcmgslvvrgaseadnrvheprtpmr 575 l:: :: :: ::: ::: Qy 532 QNMVPSSRLLVYYIVTGEQTAELVSDSVWLMIEEKCGNQLQVHLSPDADAYSPGQTVS 589</pre> | <pre>Db 576 lhiegdhkahvglvavdkavyvlnkn-kltqskvwdtvensdigctpgsgrnqvgvfada 634 ::</pre> |

Jan 28 12:22 US-08-487-283A-2.pr

US-08-487-283A-2.pr

| faccession A21898 | ecule idues | <pre>#authors Hamada, J.; Cavanaugh, P.G.; Miki, K.; Nicolson, G.L. #iournal Cancer Res. (1993) 53:4418-4423</pre> | | <pre>#accession A54561 ##molecule_type protein ##residues 25-41;749-760 ##label HAM</pre> | ##experimental_source migration-stimulating factor purified from medium conditioned by mouse hepatic sinusoidal | ENGOCHELLAL CELLS REFERENCE S16189 | #authors Sato, T.; Hong, M.H.; Jin, C.H.; Ishimi, Y.; Udagawa, N.; Shinki, T.; Abe, E.; Suda, T. | #journal FEBS Lett. (1991) 285:21-24 #title The specific production of the third component of complement | by osteoblastic cells treated with 1-alpha, 25-dihydroxyvitamin D(3). | #cross-references MUID:91293304 | Ę, | ##residues 25-31 ##label SAT | #accession SI6189 | le_type | ##residues 671-677,'X', 679-680 ##label SA2 | KEFEKENCE 149303 #authors Fev. G.; Domdev, H.; Wiebauer, K.; Whitehead, A.S.; Odink, K. | Springer Semin. Immunopathol. (1983) 6:119-147 | #title Structure and expression of the C3 gene. #cross-references MIID:84045280 | #accession 149563 | ##status preliminary ##mologija tupo mpNl | ##Wiresidus 7-5-136, Q', 138-240 ##label FEY ##wared and and and and and and and and and an | ##CIOSS-references GB:R33033; NID:q192280; CD3_KID:q192201 REFERENCE 149576 | <pre>#authors Fey, G.H.; Wiebauer, K.; Domdey, H. #iournal Ann. N. Y. Acad. Sci. (1983) 421:307-312</pre> | | nucleotide sequences of cloned cDNA. | #titles retriences notifications #accession 149576 | <pre>##status preliminary; translated from CB/EMBL/DDBJ ##malacule ture mBNa</pre> | ###################################### | COMMENT Complement C3 contains two chains, formed by removal of four | residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and | alternative complement pathways, releases the C3a anaphylatoxin | irom the amino end of the alpha chain and generates USD, which associates with the Bb fraqment of complement factor B to form | the alternative-complement-pathway C3/C5 convertage. | | COMMENT C3b, with its highly reactive thiol group, binds to the surface of | complement C5 and renders it susceptible to proteolysis by the | classical-complement-pathway C3/C5 convertase. The activity of |
|---|----------------|--|---|---|---|---|---|--|--|---------------------------------|--|---|-------------------|-------------------------------------|---|--|--|--|-------------------|--|---|--|---|---------|--------------------------------------|--|--|--|--|---|---|--|--|--|--|---|--|
| Qy 1516 IQKVCEGAACKCVEADCGQMQEELDLTISAETRKQTACKPEIAYAYKVSTTSITVENVFV 1575 | | Qy 1576 KYKATLLDIYKTGEAVAEKDSEITFIKKVTCTNA-ELVKGRQYLIMGKEALQIKYNFSFR 1634 | Db 1607 fsyliskdtwleawpleescqdadlqplcqdftefsdnlv1fgc 1650 | ۲ .ا | C3MS #type complete complement C3 precursor - mouse | CONTAINS alternative-complement-pathway U3/U3 convertase (EU 3.4,21.47) C3b subunit; C3a anaphylatoxin | ORCANISM #formal name Mus musculus #common name house mouse 30-Jun-1988 #sequence revision 30-Jun-1988 #text change | SSIONS | S16189; 149563; 149576; A01261; A05290; A29033 REFERENCE A92459 | rs | Structure of murine complement component C3: | sequence of cloned complementary and genomic DNA coding for | the beta chain. | *CLOSS-Teletenences multi-old-stold | type | ##residues 1-/24 ##label LUI #accession B92459 | ule. | ##residues 1-124 ##label LU2 grprppnrF | r.s | B.F.; Fey, G.H. | Structure of murine complement component C3: 1 | sequence of cloned complementary UNA coding for the alpha chain. | fcross-references MUID:85054819 | le type | ##residues 671-1663 ##label WET | I.B | K.; Fey, G. | | #cross-references MUID:83117730 | #contents C3a #accession A93938 | Ę, | ##residues 6/1-/48 ##label DOM REFERRICE A21898 | rs | F. B.; JOHES, C. B.; MIEKINIKKI, D. B.; Madjiussui, J.; Domuey, H.; Wetsel, R.A.; Lundwall, A.; Tack, B.F.; Fey, G.H. | # journal Proc. Natl. Acad. Sci. U.S.A. (1985) 82:9-13 | *title communi evolutionaly origin of alphaz-macrogrouniii and complement components C3 and C4. | €cross-references MUID:85113177 |

Jan 28 12:22

| (3p | City is remulated by protective cleavage involving factors H and | f | 265 n | 265 nv-data |
|---|---|----------------|--------------------|---------------------|
| I. II | I. Its degradation products can also be biologically active. | 3 . | | ; ; : — |
| COMMENT The ma | The major site of synthesis of this plasma protein is the liver. | δò | 259 V | 259 VVTEADV |
| guo | 27/2; 90/3 | q | 321 v | 321 vgkslyv |
| CATION | the list of introns may be incomplete #superfamily alpha-2-macroglobulin | ò | 319 N | : 319 NNKYLYL |
| KEYWORDS | acute phase; complement alternate pathway; complement pathway; glycoprotein; hydrolase; immune response; inflammation; plasma; serine proteinase; thiolester bond | q | 381 - | |
| FEATURE | bdameic signal processes between analyses blacks of | Qy | 379 L | 379 LDQLVGG |
| 7-7 4 25-666 | #tomain sequence *status predicted *label Jis/ #product complement C3 and C3b beta chain #status nedicted #label C3Rs. | QQ | 428 k | 428 kdtlpes |
| 25-666, 671-1663 | #product | δλ | 439 A | 439 APDLPEE |
| 671-1663 | | QQ | 488 r | 488 ryytylw |
| 671-748 749-1663 | | Qy | 497 T | 497 THYNYLI |
| 946-1303 | | QQ | 548 a | adsvwd |
| 1424-1457 | fregion p | Qy | 556 3 | 556 SDSVWLN |
| 559-816, 626-661, 693-720, 694-727, 767-750, 673-1513 | | qa | u 809 | 608 nkltqsk |
| 1101-1158, | , c. | Qy | 616 A | 616 AKKPLER |
| 1389-1458, | | QQ | и 899 | 668 rrrsvqli |
| 1518-1511, 1518-1590, 1537-1661 | | Qy | 675 PI | 675 PRRTLQ-I |
| 1537-1646 1637-1646 748-749 | #disulfide_bonds #status predicted\ #r pauane site Arn-Ser (C3 convertage) #status | qq | 728 CI | 728 cnhitkly |
| 930 1617 | predicted the predicted the predict of the predicted the predicted the predicted the predict of | οy | 732 0 | cvvasqu |
| 1701 '656 | *Dinding Site carbonydrate (ABN) (covatent) *Bratus predicted\ | qq | 788 t | 788 tkvmnif |
| 1010-1013 1303-1304 | <pre>#cross-link thiolester (Cys-Cln) #status predicted\ #cleavage site Arg-Ser (complement factor I) #status</pre> | δŷ | 784 -1 | : : 784 -KQLQFA] |
| 1320-1321 | predicted february february | qq | 848 r | 848 ravlfny |
| SUMMARY | predicted flength 1663 #molecular-weight 186482 #checksum 646 | ογ | 842 K | KGTVYNYI |
| Query Match Best Local Similarity 28.28; | Score 2296; DB 2; Length 1663; Pred. No. 0.00e+00; | QQ | 901 v ₁ | 901 vplkigg : |
| Matches 479; | Conservative 447; Mismatches 665; Indels 106; Gaps 84; | δ | 901 L | 901 LPLEIGL |
| Db 28 ysiitpnv | 28 ysiitpnvlrleseetivleahdaqqdipvtvtvqdflkrqvltsektvltqasqhlrsv 87 | q ₀ | м 096 | 960 vdvp-aao |
| Qy 23 YVISAPKI | /YGYTEAFDATISIKS | Qy | 957 KI | 957 KEFPYRII |
| Ub 88 sikipask | 88 sikipaskefnsdkeghkyvtvvanfgetvvekavmv¤fgsgylfiqtdktlytpgstvl 147 : :::: : : : : | qq | 1018 mtptviav | tptvia |
| Qy 83 AILTIQPE | 3QNPVSYVYLEVVSKHFSKS | δ | 1015 VVPVFYVI | /PVFYVI |
| Db 148 yriftvdnnllp | 148 yriftvdnnllpvgk-tvvilietpdgipvkrdilssnnqhgilpls-wnipelvnmggw 205 | QQ | 1075 st | stwitay |
| Qy 143 VRVYSLNI | AKRETVLTFID-PEGSEVDMVEEI | δy | 1075 S | STWLTAF |
| Db 206 kirafyeh | 206 kirafyehapkqifsaefevkeyvlpsfevrveptetfy-yiddpnglevsiiakflygk 264 | qa | 1135 -1 | -nakead |
| Qy 200 TIKAKYKE | 200 TIKAKYKEDFSTTGTAYFEVKEYVI.PHFSVSIEPEYNFIGY-KNFKNFEITIKARYFYNK 258 | δò | 1135 VEARENS | ARENS |

18-08-487-283A-2.rpr tafvifgv-qd--gdkkislahsltrvviedgvgdavltrkvlmegvrpsnadal 320 Jan 28 12:22

| 셤 . | 265 | rrpsnad | ⊼ : |
|-----|------|--|------|
| ò | 259 | .REDIKDDQKEMMQTAMQNTMLINGIAQVTFDSETAVKELSYYSLED | 318 |
| ପ୍ର | 321 | nveaersgipivtspygihftktpkffkpampfd | 380 |
| ò | 319 | STGGFSEEAEIPGIKYVLSPYKLALVATPLFLKPGIPYPIKVQVKDS | 378 |
| g | 381 | -dgs-pas-kvl-v-vtgg-snakaltq-ddgvaklsintpnsrgpltitvrtk | 427 |
| δ | 379 | SVPVILNAQTIDVNQETSDLDPSKSVTRVDDGVASFVLNLPSGVTVLEF | 438 |
| g | 428 | kdtlpesrqatktmeahpystmhnsnnylhlsvsrmelkpgdnlnvnfhlrtdpgheaki | 487 |
| à | 439 | IIVTPKS-P-Y | 496 |
| g | 488 | ryytylvmnkgkllkagrqvrepgqdlvvlslpitpefipsfrlvayytligasggrevv | 547 |
| 0y | 497 | SRLLVYY IVTGEQTA- | 555 |
| q | 548 | vkdscigtlvvkgdprdnhlapgqqttlriegnqgarvglvavdkgvfvln | 209 |
| 0y | 556 | MINIEEKCGNQLQVHLSPDADAYSPGQTVSLNMATGMDSWVALAA | 615 |
| qq | 809 | nkltqskiwdvvekadigctpgsgknyagvfmdaglafktsgglqtegradlectkpaar | 199 |
| δ | 616 | PLERVFQF1EKSDLGCGAGGGLNNANVFHLAGLTFLTNANADDSQENDEPC-KEI | 674 |
| q | 899 | merrmdkaggytdkglrkccedgmrdipmryscgrrarlitggencikafid | 727 |
| δλ | 675 | PRRTIQ-K-KIEELAAKYKHSVVKKCCYDGAC-VNNDETCEQRAARISLGPRCIKAFTEC | 731 |
| q | 728 | cnhitklreghrrdhvlglarseleediipeediisrshfpgswlwtieelkepekngis | 787 |
| δý | 732 | OLRANISHKD-MOLGRLHMK-TLLPVSKPEIRSYFPESWLMEVH-L- | 783 |
| qq | 788 | tkvmniflkdsittweilavslsdkkgicvadpyeirvmqdffidlrlpysvvrneqvei | 847 |
| δ | 784 | .ALPDSLTTWEIGGIGISNT-GICVADTVKAKVFKDVFLEMIPYSVVRGEQIO | 841 |
| QQ | 848 | bafcsmataknryfqtikippkssvavpyvi | 900 |
| δλ | 842 | CVKMSAVEGICTSESPVIDHQGTKSSKCVRQ-KVEGSSSHLVTFTV | 900 |
| q | 901 | ggqevevkaavfnhfisdgvkktlkvvpegmrinktvaih | 959 |
| ý | 901 | EIGLHNINFSLETWFGKEILVKTLRVVPEGVKRESYSGV-TLDPRGI-YGTISRR | 926 |
| QQ | 096 | vdvp-aadlsdqvpdtdsetrii-lqgspvvqmaedavdgerlkhlivtpagcgeqnmig | 1017 |
| δλ | 957 | PYRIPL-DLVPKTEI-KRILSVKGLLVGEILSAVISQEGINILTHLKGSAEAELMS | 1014 |
| QQ | 1018 | ldqteqwekfgiek-rq-ealel-ikkgytqqlafkqpssay | 1074 |
| ٥y | 1015 | YVFHYLETGNHWNIFHSDPLIEKQKLKKKLKEGMLSIMSYRNADYSYSY | 1074 |
| QQ | 1075 | Layvvkvfslaanliaidshvlcgavkwlilekqkpdgvfqedgpvihqemiggfr | 1134 |
| ò | 1075 | THILTAFALRVI.GOVNKTVEDNONS I CNSILLMLVENYQLDNGSFKENSQYQE | 1134 |
| g | 1135 | -nakeadvsltafvlialqeardicegqvnslpgsinkageyieasymnlqrpytvaiag | 1193 |
| δ | 1135 | nslyltaetvigirkaedicp-luk-idtalikadnellentipagstetlaisa | 1192 |

!product C3a anaphylatoxin #status predicted #label C3T\

*product complement C3 alpha chain #status predicted

#label CC3A\

product complement C3b alpha' chain #status predicted

product C3d fragment #status predicted #label C3D\

!region properdin binding\

1007-1308 1429-1461

Amino acid sequence of guinea pig C3a anaphylatoxin

cross-references MUID:89113342

##molecule_type protein

accession

##residues

Gerard, N.P.; Lively, M.O.; Gerard, C. Protein Seq. Data Anal. (1988) 1:473-478

1-1666 ##label AUE

##molecule_type mRNA A37156

accession

*residues

#fcross-references GB:M34054

503375

Pauthors journal

REFERENCE

title

##experimental_source complement-activated guinea pig serum

676-730,'N',732-752 ##label GER

951-1308

product C3dk fragment #status predicted #label

flabel

154-1666

676-753

1-22 23-671

ftitle Molecular basis of complement C3 deficiency in guinea pigs. fcross-references MUID:90307998

Auerbach, H.S.; Burger, R.; Dodds, A.; Colten, H.R. J. Clin. Invest. (1990) 86:96-106

#authors | journal

ACCESSIONS

DATE

REFERENCE

A37156; S03375; A20342; D20342; C20342; A31222

16-Feb-1997

#formal name Cavia porcellus #common name guinea pig 07-Feb-1992 #sequence_revision 07-0ct-1994 #text_change

alternative-complement-pathway C3/C5 convertase (EC

CONTAINS ORGANISM

TITLE ENTRY

complement C3 precursor - guinea pig

#type complete

8

3.4.21.47) C3b subunit; C3a anaphylatoxin

FEATURE

KEYWORDS

COMMENT

NH2-terminal structure and cleavage of guinea pig pro-C3, the precursor of the third complement component. C3b is regulated by proteolytic cleavage involving factors H and I. Its degradation products can also be biologically active. C3b, with its highly reactive thioi group, binds to the surface of inflammation; liver; plasma; serine proteinase; thiolester Complement C3 contains two chains, formed by removal of four residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and alternative complement pathways, releases the C3a anaphylatoxin from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the #domain signal sequence #status predicted #label SIGN #product complement C3 and C3b beta chain #status predicted #label C3BBN product complement C3 #status predicted #label CC3\
iproduct complement C3b #status predicted #label C3B\ The major site of synthesis of this plasma protein is the liver. classical-complement-pathway C3/C5 convertase. The activity of Identification and alignment of a thiol ester site in the Goldberger, G.; Thomae, M.L.; Tack, B.F.; Williams, J.; Colten, H.R.; Abraham, G.N. J. Biol. Chem. (1981) 256:12617-12619 ##molecule_type protein #fresidues 993-1012,1014-1017,'E',1019-1030,'Y' ##label TH2 C3a anaphylatoxin is a vasoactive peptide and a mediator of inflammation. acute phase; complement alternate pathway; complement pathway; glycoprotein; hydrolase; immune response; the alternative-complement-pathway C3/C5 convertase. missing the carboxyl-terminal arginine third component of guinea pig complement. #superfamily alpha-2-macroglobulin Thomas, M.L.; Tack, B.F. Biochemistry (1983) 22:942-947 676-687 ##label TH1 23-38 ##label GOL A20342 D20342 A20342 C20342 23-671, 676-1666 23-671, 754-1666 676-1666 CLASSIFICATION

COMMENT COMMENT

1603 KVTCTNA-ELVKGRQYLIMGKEALQIKYNFSFRYIYPLDSLTWIEYWPRDTTC-S-SCQA 1659

form isolated is inactive C3a anaphylatoxin and US-08-487-283A-2.pg cross-references MUID:83178889 #cross-references MUID:82075767 ##molecule_type protein ##molecule_type protein ##residues ##residues #accession #accession #accession Jan 28 12 22 ##note #authors # journal authors journal title REFERENCE #title REFERENCE COMMENT S 1295 dvsfhlpsrssattfrllwengnllrseetkqneafsltak-gkgrgtlsvvavyhaklk 1353 1409 tgfapdtkdlellasgvdryiskyemnkafsnkntliiylekishteedcltfkvhgyfn 1468 1194 yalalmnkleepy--lgkflnt-a--k-drn--r-w-e--ep-dqql-yn----veats 1234 1193 YALSLGDKTHPQFRSIVSALKREALVKGNPPIYRFWKDNLQHKDSSVPNTGTARMVETTA 1252 1235 yallalllikdfdsvppvvrwlneqryygggygstqatfmvfqalaqyqtdvpdhkdlnm 1294 1312 DIDVSYKHKGALHNYKMT-DKNFLGRPVEVLINDDLIVSTGFGSGLATVHVTTVVHKTST 1370 1354 skvtckkfdlrvsirpapetakkpeeakntmfleictky-1g-dvdat-ms--ildismm 1408 1469 vgliqpgsvkvysyynleesctrfyhpekddgmlsklchsemcrcaeencf-mqqsqe-k 1526 1527 inInvrldkacepgvdyvykteltnikllddfdeytmtiqqviksgsdevqagqrkfis 1586 1587 hikcrnalklqkgkkylmwglss-dl-wgekpntsyiigkdtwvehwpeaeecqdqkyqk 1644

::

_ :: ::

<u>-</u>

셤

ð

유

Š

셤

ð g ð

Š

a

ð g δ g ð 셤

23

US-08-487-283A-2.pr

| 1394-1462, 1510-1515. | | qΩ | 663 gldcpl |
|---------------------------------|--|------|----------------|
| 1522-1593, 1540-1664. | | ٥y | 666 DEPC-F |
| 1640–1649 753–754 | <pre>#disulfide bonds #status predicted\ #cleavage_site Arg-Ser (C3 convertase) #status</pre> | QQ | 723 eacvka |
| 944, 1620 | <pre>predicted\ #binding site carbohydrate (Asn) (covalent) #status</pre> | y g | 722 PRCIKA |
| 1015-1018 1308-1309 | <pre>#cross-link thiolester (Cys-Gln) #status experimental/ #cleavage site Arg-Ser (complement factor I) #status</pre> | g Vy | 179 L-VP-F |
| 1325-1326 | predicted predicted formula fo | QQ | 843 svvrne |
| SUMMARY | predicted #length 1666 #molecular-weight 186487 #checksum 5419 | Qy | 832 SVVRGE |
| Query Match Rest Local Simil | n 19.1%; Score 2302; DB 2; Length 1666; Similarity 28.5%: Pred: No. 0.00e+00: | Ob | 897 ssvavp |
| Matches 486; | 486; Conservative 450; Mismatches 658; Indels 111; Gaps 83; | ٥y | 892 SSHLVT |
| Db 23 dpmysiit | 23 dpmysiitpnilrleneetvvleahevqqdipvtvtvhdfpakknvlssektvltsatgy 82 | qq | 957 lgqggv |
| Oy 20 EQTYVISH | 20 EQTYVISAPKIFRVGASENIVIQVYGYTEAFDATISHI3YPDKKFSYSSGHVHLSSENKF 79 | Qy | 949 I-YGTI |
| Db 83 lgtvtik | 83 lgtvtikipaskefksdkgr-klvvvqaafggtqlekvvlvslqsgylfiqtdktiytpg 141 : : { : : : : | Ωρ | 1014 gcgeqn |
| Qy 80 QNSAILT | 80 QNSAILTIQP-KQLPGGQNPVSYVYLEVVSKHFSKSKRMFITYDNGFLFIHTDKPVYTPD 138 | Qy | 1006 GSAEAE |
| Db 142 stvlyrii | 142 stvlyriftvdsdllpvgrtiivtietpdgipikrdtlssnnqhgilpls-wnipelvnm 200 | QΩ | 1071 yaafkn |
| Qy 139 QSVKVRV) | 139 QSVKVRVYSLNDDLKPAKRETVLIFIDPEGSEVDMVEEIDHIGIISFPDFKIPSNFRY 196 | ٥y | 1066 YSVWKG |
| Db 201 gqwkiqa1 | gqwkiqafyenspkqvfsaefevkeyvlpsfevlveptekfy-yiddpkglevniiarf1 259 | ηρ | 1131 hqemig |
| Qy 197 GMWTIKA | 197 GMHTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFIGY-KNFKNFEITIKARYF 255 | 0y | 1126 PIKLQG |
| 0b 260 ygknv-do | 260 ygknv-dgtafvifgv-qdgdgrislagsltrvviedgsgevvlsrqvlldgvgpsrp 315 | qu | 1190 rpytla |
| Qy 256 YNKVVTEZ | 256 YNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLINGIAQVTFDSETAVKELSYYSL 315 | 6y | 1184 STFTLA |
| Db 316 ealvgks] | 316 ealvgkslyvsvtvilhsgsdmveaersgipivtspygihftktpkyfkpampfeimvlv 375 | qa | 1235ve |
| Qy 316 EDLANKYI | TXIAVTVIESTGGFSEEAEIPGIKYVLSPYKLNLVATPLFIKPGIPYPIKVQV 375 | Qy | 1244 TARMVE |
| 0b 376 tnp-d | 376 tnp-dgs-pap-hvp-v-vtgg-snvgsltgad-gvarlsintpntrgplsvtv 422 | Ob | 1291 vpdhkd |
| Qy 376 KDSLDQLV | 376 KDSLDQLVGGVPVILMAQTIDVNQETSDLDPSKSVTRVDDGVASFVLMLPSGVTVLEFNV 435 | Qy | 1304 VKQLR- |
| Db 423 qtkkggip | 423 qtkkggipdarqaintmqalpyttmynsnnylhlsmprtelkpgetinnnfhlrsdpnqe 482 | QQ | 1350 vavyya |
| Qy 436 KTDAPDLE | 436 KTDAPDIPEENQARECYRAIAYSSISQSYLYIDMTDN:XALLVGEHINIIVTPKS-PYID 494 | ۵y | 1362 TTVVHK |
| Db 483 akiryyty | 483 akiryytylimnkgkllkvgrqprepgqalvvlpmpitkelipsfrlvayytligasaqr 542 | QD | 1407 11di |
| Qy 495 -KITHYNY | GTREKFSDASYQS | θŷ | 1421 HAVMDI |
| Db 543 evvadsva | evvadavwadvrdscvgilvvkggsgkdgqdkrqqhlprqqmtlriegnqgarvglvavd 602 : : :: ; | qq | 1465 fkihqf |
| Qy 553 ELVSDSVN | ELVSDSVWINIEEKCGNQLQVHLSPDA-DAYSPGQTVSIAMATGMDSWVAIAAVD 606 | δy | 1477 FRIFEL |
| Db 603 kgvfvlnk : ::: | 603 kgvfvlnkkhkltgsklwdvvekadigctpgsgkdyagvftdaglsfksskaglqtaqre 662 : : : | qu | 1524 iqlp : |
| Qy 607 SAVYGVQF | rgakkplervpofleksdlgcgaggelnnanvfhlagltfltn-anaddsoen 665 | Qy | 1535 MQEELD |

| đ | 663 | gldcpkpaarrrrsvqlmerrmdkagkykskelrrccedgmrenpmqfscgrraryvsig 722 |
|-----|------|--|
| ολ | 999 | |
| đ | 723 | eacvkafldcctymaqlrqqhrreqnlqlarsdmdediipeediisrsqfpsswlwtiee 782 : : : : :: : :: |
| ò | 722 | PRCIKAFTECCVVASQLRA-NISHKDMQLGRLHMK-TLLPVSKPEIRSYFPESWLWEVH- 778 |
| qq | 6 | erngietktmniflkdsittweilavsledkkgicvadpfevtvmqdffidlrlpy |
| 0y | 779 | LTTWEIQGIGISNT-GICVADTVKAKVFKDVFLEMNIPY 83 |
| q | 843 | svvrneqveiravlynyreaq-slkvr-vellhnpafcslatakkrhtqtvtigpk 896 |
| δ | 832 | SVVRGEQIQLKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKSSKCVRQKVEGS 891 |
| qq | 897 | ssvavpyulvplkiglqevevkaavynyfisdgvkkllkvvpegmrunktvairtlnpeq 956 |
| δλ | 892 | SSHLVTFTVLPLEIGLHNINFSLETWFGKEILVKTLRVVPEGVKRESYSGV-TLDPRG 948 |
| qq | 957 | lggggv-greeip-aadlsdgvpdtdsetkil-lggtpvagmaedavdaerlkhliitps 1013 |
| ολ | 949 | -YGTIS |
| Ор | 1014 | gcgeqnmigmtptviavhyldqteqwekf-glekrqealnlinrgytqqlafkqpnwa 1070 |
| Qy | 1006 | AELMSVVPVFYVFHYLETGNHMVIFHSDPLIEKQKLKKKLKEGMLSIMSYRN |
| QQ | 101 | yaafknrasstwltayvvkvfslaanligidsevlcgavkwlilekqkpdgvfqedgpvi 1130 |
| δ | 1066 | YSVWKGCSASTWLTAFALRVLGQVNKTVEQNQNSICNSLLMLVENYQLDNGSFKENSQYQ 1125 |
| qq | 1131 | hqemiggvrt-aqeadvsltafvlialqeakdicraqvnnleaninkagdyiesryadvr 1189 |
| δý | 1126 | PIKLOGTLEVEARENSLYLTAFTVIGIRKAFDIG-PLVK-IDTALIKADNFLLENTLPAQ 1183 |
| QQ | 1190 | rpytlaiagyalallerlngatlqk-fl-naateknr-weea-rqkl-ys- 1234 |
| δy | 1184 | STFTLAISAYALSLGDKTHPQFRSIVSALKREALVKGNPPIYRFWKDNLQHKDSSVPNTG 1243 |
| ΩP | 1235 | veatsyallalllkdfdavppvvrwlnegryygrgygstgatfmvfgalagygtd 1290 |
| Ø | 1244 | TARMVETTAYALLTSINIKDINYVNPVIKWLSEERYGGGFYSTODTINAIEGLTEYSLL 1303 |
| qq | 1291 | hkdlnmevalqlpsrsspskfrlvweagsllrseatkqnegfklt. |
| δy | 1304 | DVSYKHKGALHNYKMT-DKNFLGRPVEVLLNDDLIVSTGFGSGLATVH |
| qq | 1350 | vyyaktkrkvvcknfdlrvtlkpapdtvkkpqeakstmilgictry-lg-dqdat-m |
| ۵y | 1362 | TTVVHKTSTSEEVC-SFYLKIDTQDIEASHYRGYGNSDYKRIVACASYKPSREESSSGSS 1420 |
| qq | 1407 | ildismmtgfipdtddlkllatgvdryiskyemnkdfskntliiyldkvshseecls 1464 |
| δλ | 1421 | HAVMDISIPTGISANEEDIKALVEGVDQLFTDYQI-KD-G-H-VILQLNSIPSSDFLCVR 1476 |
| QQ | 1465 | fkihqffnygliqpgsvkvysyynldetctqfyhpekedgmlnklchkdlcrcaeencf- 1523 |
| δλ | 1477 | FRIFELFEVGFLSPATFTVYEYHRPDKQCTMFY-S-TSNIKIQKVCEGAACKCVEADCGQ 1534 |
| qq | 1524 | iqlpekitlderlekacepgvdyvyktkllkmeleddfdeyimtieqvikagsdevqa 1581 : :- - |
| ٥'n | 1535 | KPEIAYAYKVSITSITVENVFVKYKAT |

ð

8

US-08-487-283A-2.pg

25-666, 671-1663 25-666, 749-1663 671-1663 671-748 REFERENCE title REFERENCE 52-666 #title KEYWORDS COMMENT COMMENT COMMENT COMMENT FEATURE Jacobs, J.W.; Rubin, J.S.; Hugli, T.E.; Bogardt, R.A.; Mariz, I.K.; Daniels, J.S.; Daughaday, W.H.; Bradshaw, R.A. Biochemistry (1978) 17:5031-5038 Nucleotide and deduced amino acid sequence of rat complement Suzuki, Y.; Chang, H.W.; Hara, S.; Inoue, K. Proc. Natl. Acad. Sci. U.S.A. (1990) 87:2395-2399 Proteinaceous inhibitors of phospholipase A-2 purified from 1582 gkerrfishikcrdalhlkegkhylmwglss-dl-wgerpnmsyiigkdtwveawpeaee 1639 1595 DSEITFIKKVTCTNA-ELVKGRQYLIMGKEALQIKYNFSFRYIYPLDSLTWIEYWPRDTT 1653 Suwa, Y.; Kudo, I.; Imaizumi, A.; Okada, M.; Kamimura, T.; alternative-complement-pathway C3/C5 convertase (EC 3.4.21.47) C3b subunit; C3a anaphylatoxin #formal_name Rattus norvegicus #common_name Norway rat 04-Dec-1992 #sequence_revision 07-Oct-1994 #text_change #title Purification, characterization, and amino acid sequence
 rat anaphylatoxin (C3a).
#cross-references MUID:79062262
#accession A01260 S15764; A54562; A01260; B35979; A35979; PN0567; PN0566; Sundstrom, S.A.; Komm, B.S.; Ponce-de-Leon, H.; Yi, Z.; Teuscher, C.; Lyttle, C.R. J. Biol. Chem. (1989) 264:16941-16947 Estrogen regulation of tissue-specific expression of 671-703,'K',705-720,'KL',723-748 ##label JAC C3RT #type complete complement C3 precursor - rat 37K phospholipase A2 inhibitory protein three disulfide bonds are present Misumi, Y.; Sohda, M.; Ikehara, Y. Nucleic Acids Res. (1990) 18:2178 ##molecule_type mRNA
#fresidues 'P',1316-1595 #flabel SUN ##molecule_type protein ##residues 'X', 998-1005 ##label SUW inflammatory sites in rats. translation not shown ##molecule_type_mRNA #fresidues 1-1663 ##label MIS ##cross-references EMBL:X52477 1654 C-SS-CQAFLANLDEFAEDIFLNGC 1676 1640 cqdeenqqqcqdlgtftenmvvfgc 1664 <u>:</u> ##cross-references GB:M29866 fcross-references MUID:90245672 \$cross-references MUID:90207203
\$accession B35979 A32281; S08692 S15764 complement C3. 16-Feb-1997 ##molecule_type protein \$15764 A54562 A35979 A35979 A54562 g ##residues #residues ALTERNATE NAMES ##status ##status *accession #accession *accession ##note #authors #authors # journal # journal #authors #authors # journal f journal ACCESSIONS ftitle #title title REFERENCE REFERENCE REFERENCE REFERENCE CONTAINS ORGANISM 쇰

##molecule_type protein ##residues 25-41 ##label KUI ##experimental_source 17beta-estradiol-stimulated uterus of immature rat Kuivanen, P.C.; Capulong, R.B.; Harkins, R.N.; DeSombre, E.R. C3b, with its highly reactive thiol group, binds to the surface of foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the classical-complement-pathway C3/C5 convertase. The activity of C3b is regulated by proteolytic cleavage involving factors H and I. Its degradation products can also be biologically active. The major site of synthesis of this plasma protein is the liver. distinct, homologous protein rather than as an alternatively processed form or fragment Complement C3 contains two chains, formed by removal of four residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and amino end of peptide designated neutrophil chemotactic factor 1 and probably identical to C3a anaphylatoxin alternative complement pathways, releases the C3a anaphylatoxin proteins are structurally related to complement component Complement component C3-derived neutrophil chemotactic factors purified from exudate of rat carrageenin-induced Biochem. Biophys. Res. Commun. (1989) 158:898-905 The estrogen-responsive 110K and 74K rat uterine secretory from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form response; inflammation; liver; plasma; serine proteinase; #product complement C3 and C3b beta chain fstatus
 predicted flabel C3BN
product complement C3 fstatus predicted flabel CG3\
 fproduct complement C3 fstatus predicted flabel C3B\
 fproduct complement C3 alpha chain fstatus predicted
 fproduct complement C3 alpha chain fstatus predicted the authors treat this 74K uterine secretory protein, identical as far as sequenced to complement C3, as domain signal sequence #status predicted #label SIG\ amino end of a C3-derived peptide designated exudate neutrophil chemotactic factor 2, or C3-beta-c C3a anaphylatoxin is a vasoactive peptide and a mediator of acute phase; chemotaxis; complement alternate pathway; complement pathway; glycoprotein; hydrolase; immune Biochem. Biophys. Res. Commun. (1993) 194:1181-1187 the alternative-complement-pathway C3/C5 convertase, #superfamily alpha-2-macroglobulin Nakagawa, H.; Komorita, N. ##molecule_type protein ##Label NA2 ##residues 671-687 ##Label NA2 568-592 ##label NAK thiolester bond #cross-references MUID:89149812 inflammation. ##molecule_type protein inflammation. PN0566 PN0566 PN0567 ##residues CLASSIFICATION faccession | faccession #accession ##note ##note authors f journal #authors journal j

380 -d---gs-par-rvp-v-vtqg-sd---aqaltq-ddgvaklsvntpnnrqpltitvstk 426

õ g ð ď 8 임 ð

427 kegipdarqatrtmqaqpystmhnsnnylhlsvsrvelkpgdnlnvnfhlrtdagqeaki 486

487 ryytylvmnkgkllkagrqvrepgqdlvvlslpitpefipsfrlvayytliganggrevv 546

497 THYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPSSRLLVYYIVTGEQTA-ELV 555

US-08-487-283A-2.rpi Jan 28 12:22

33

| OP | 547 | adsvwvdvkdscvgtlvvkgdprdnrqpapghqttlriegnqgarvglvavdkgvfvlnk | 909 |
|----------|------|--|------|
| οy | 556 | : ::: : : : : : : | 614 |
| QQ | 607 | knkltgskiwdvvekadigctpgsgknyagvfmdagltfktngglqtdgredpecakpaa | 999 |
| δý | 615 | GARKP LERVFQF LEKSDLGCGAGGGIANANVFHLAGLTFLTNANADDSQENDEPC-KEIL | 673 |
| Op | 199 | rrrrswglmerrmdkaggytdkglrkccedgmrdipmpyscgrrarlitggesclkafmd | 726 |
| ò | 674 | RPRRTLQ-K-KIEEIAAKYKHSVVKKCCYDGAC-VNNDETCEQRAARISLGPRCIKAFTE | 730 |
| 8 | 727 | ccnyitklreqhrrdhvlglarsdvdediipeediisrshfpeswlwtieelkepekngi | 786 |
| à | 731 | CCVVASQLRANISHKD-MQLGRLHMK-TLLPVSKPEIRSYFPESWIMEVH-L-VP-RR- | 783 |
| DP | 787 | | 846 |
| ý | 784 | ∃ ≲ | 840 |
| g | 847 | iravlfnyreqeklkvr-vellhnpafcsmatakkryyqti-e-ippkssvavpyvi | 900 |
| Š | 841 | IKGTVYNYRTSCMQFCVKMSAVECICTSESPVIDHQGTKSSKCVRQKVEGSSSHLVTFTV | 900 |
| QQ | 901 | vplkiglqevevkaavfnhfisdgvkkilkvvpegmrvnktvavrtldpehlngggvqre | 960 |
| ò | 901 | LPLEIGLHNINFSLETW-FGKEILVKTLRVVPEGVKRESYSGV-TLDPRGIY-GTISRR | 956 |
| g | 961 | dvnaadls-dqvpdtdsetril-lqgtpvaqmaedavdgerlkhlivtpsgcgeqnmigm | 101 |
| ζý | 957 | KEFPYRIPLDIAPKTEI-KRILSVKGLIVGEILSAVLSQEGINILTHIPKGSAEAELMSV | 101 |
| Db | 1019 | tptviavhyldqteqwekf-glekrqealelikkgytqqlafkqpisayaafnnrpps | 107 |
| δ | 1016 | THILT CHIMNIEHSDPLIEKQKIKKKIKEGMLSIMSYRNADYSYSVW | 107 |
| PP PP | 1076 | twltamwsrsfslaanliaidsqvlcgavkwlilekqkpdgvfqedgpvihqemiggfr- | 1134 |
| õ | 1076 | THILL THE | 113 |
| ДQ | 1135 | ntkeadvsltafvlialqeardicegqvnslpgsinkageyleasylnlqrpytvaiagy | 1194 |
| à | 1136 | SLYLTAFTVIGIRKAFDICP-LVK-IDTALIKADNFLLENTLPAQSTFTLAI | 119 |
| Op | 1195 | alalmnkleepyltkflnt-a-k-drnr-weepggql-ynveatsy | 123 |
| ζ | 1194 | LVKGNPPIYRE | 125 |
| q | 1236 | allalllikdfdsvppvvrwlnderyygggygstgatfmvfgalagyradypdhkdlnmd | 129 |
| δ | 1254 | ALTSINIKDINYVNPVIKWISEEQRYGGGFYSTQDTINAIEGLIEYSLLVKQLR-LSMD | 1312 |
| g Q | 1296 | vslhlpsrssptvfrllwesgsllrseetkqnegfsltak-gkqqqllsvvtvyhakvkg | 1354 |
| ð | 1313 | VEVLLNDDLIVSTGFGSGLAT | 1371 |
| g | 1355 | kttckkfdlrvtikpapetakkpqdakssmildictry-lg-dvdat-msildismmt | 140 |
| à | 1372 | EEVCS-FYLKIDIQDIEASHYRGYGNSDYKRIVACASYKPSREESSSGSSHAVMDISLPT | 143(|
| ą | 1410 | gfipdtndlellssgvdryiskyemdkafsnkntliiylekishseedclsfkvhqffnv | 1469 |
| ð | 1431 | PSSDFLCVRFRIFE | 1485 |

350 tpkyfkpgmpyeltvyvtnp-d---gs-paa-hvp-v-vs-ea--fh-smgtt-lsdgta 396

셤

US-08-487-283A-2.mr Jan 28 12:22

8

1037 aqqmvykkadhsyaaftnrassswltayvvkvfamaakmvagisheiicggvrwlilnrq 1096 LSIMSYRNADYSYSVWKGGSASTWLTAFALRVLGQVNKYVEQ-NQNSICNSLLMLVENYQ 1112 qpdgafkenapvlsgtmqgg--iqgaeeevyltafilvalleskticndyvnsldssikk 1154 atnyllkkyeklgrpyttaltayalaaadqlndd-r-v--lm-aa-stgrd---h-w-e 1202 | | :|| : | :| |:|||: | : | | | : | 1203 --ey-na---ht---hniegtsyallallkmkkfdqtgpivrwltdqnfygetygqtqat 1253 skinhliitpsgcgeqnmirmaapviaty-yldtteqwetlginrrte-a-vnqivt-gy 1036 1171 ADNFLLENTLPAQSTFTLAISAYALSLGDKTHPQFRSIVSALKREALVKGNPPIYRFWKD 1230 1254 vmafqalaeyeiqmpthkdlnlditielpdrevpiryrinyenallartvetklnqditv 1313 959 111 169 820 kvffidlqmpysvvkneqveirailhnyvnedi-y-vr---velly---npafcsastkg 863 980 359 TPLFLKPGIPYPIKVQVKDSLDQLVGGVPVILNAQTIDVNQETSDLDPSKSVTRVDDGVA 418 397 klilniplnaqslpitvrtnhgdlprerqatksmtaiayqtqggsgnylhvaitsteikp 456 571 596 631 691 751 880 994 KDVFLEMNIPYSVVRGEQIQIKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKS RESYSGYTLDPRGIYGTISRRKE-FPYRIPLDLVPKTEIKRILSVKGLLVGEILSAVLSQ gdnlpvnfnvkgnanslkqikyftylilnkgkifkvgrqprrdgqnlvtmnlhitpdlip |::| :: | :: : | : |||||||||| | : : ::|: :| ::|
GEHINI-1-VTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVP sfrfvayyqv-gnn--eivadsvwvdvkdtcmgtl-v-vkgdnliqmpgaamkiklegdp ANADDSQENDEPCKE-ILRPRRT---LOKKIEEIAAKYKHSVVKKCCYDGACVNNDE-TC garvglvavdkavyvlndkykisgakiwdtieksdfgctagsggnnlgvfedaglaltts tnlntkgrsaakcpqpanrrrrssvllldsnaskaaefqdqdlrkccedvmhenpmgytc ekrakyiqegdackaafleccryikgvrdenqreselflarddnedgfiadsdiisrsdf ksivtivkldprak-gvggtqlevikarklddrvpdteietkiiiqgdpvaqiiensidg _ :: 1054 1097 1155 457 479 537 597 657 692 712 752 770 812 821 864 881 936 517 572 632 922 981 g a g g g 유 유 유 음 염 셤 유 Š ð ò ð ð g g g g ð Š ò ò ò ò ð ò Š Š Š

ð අ ò 셤 à 9 ð D P õ g Š

US-08-487-283A-2.pr

| 902 937 960 997 1017 1056 1116 1116 | ##molecule_type mRNA #fresidues |
|---|---|
| | #cross-references MID:92192016 #accession S21045 #fetatus preliminary #fetatus preliminary #fetasidus |
| Db 790 fkqfflkvhtpyalkqyeqvelrvviynymnqdvkgeiqvkcgdgict-daeqneplk : :: :: :: : : : : : : :: | ss a comple |
| Db 730 rshfpesfmweiiklsraaengksritkkmpdsittwdiqavevsgskglcvgpsleltv 789 | |
| Db 670 lsceertkhihdegegcgetflecckhveeelliameeededlgrsggedfmiqesgvvi : : : | 521045 |
| Db 611 aimakeisgmddvkdpgcpnghtrrkrelvleiaiekasty-paelrkccrdaaiesplr 669 ::::: : : : : : :: : | 1637 YPLDSLTWIEYWPRDTTCSSCQAFLANL-DEFAE 1669 |
| Db 551 isgesdswvavgvvdkaayvldkknkltankvykameasdlgcsvgsgktgplvfrdag1 610 :: | 1580 TLIDIYKTGEAVAEKDSEITFIKKVTCTNA-ELVKGRQYLIMG-K-EALQIKYNFSFRYI 1636 1601 it-kn-twierwphedecqe-eef-qklcddfaq 1630 |
| Db 494 pelvpqfrlvaffilpsgelvadsiiidvkdschaklsldvaggkrlfsprdnvnfd :: : :: :: : : : : : | CEGAACKCVEADCGOMQEELDLTISAETRKQTACKPEIAYAYKVSITSITVENVFVKYKA 1579 dvlevikqgtdenprakthqyisqrkcqealnlkvnddyliwqsrsdllptk-d-kisyi 1600 |
| Qy 472 NHKALLVGEHLANIVTPKSPYIDKITHYNYLLLSKGKIIHFGTREKFSDASYQSINIPVT | cignvcrcagetcsslnhq-e-ridvplqiekacetnvdyvyktkllrieeqdgndiyvm 1542 |
| | 1422 lyinkvsneedecinikilknievyjiqqgavkvysyynidekcikrynpdkytqilnki 1484 : :: :: |
| Db 379 -mdngiamqtintarnsklnikvqtederleqsqqaeasftiasysspsgsfihln-a-: : : : : :::: Qy 412 RVDDGVASFVLNLPSGVTVLEFNVKTDAPDLPEENQAREGYRALAYSSLSQSYLYLDWTD | SYKPSREESSSGSSHAVMDISLPTGISANEEDIKALVEGVDQIFTDYQI-KD-GHVI 1461 ivlnkvshsedeclhfkilkhfevqfiqqqsvkvysvynldekctkfyhpdkqtqllnki 1484 |
| Qy 352 YKLMLVATPLETKPGIPYPIKVQVKDSLDQLVGGVPVILMAQTIDVNQETSDLDPSKSVT | 1370 ry-1gevds-tm-tiidism tgf1pdaed tr1skgvdryisryevdnnmaqkvavi 1424 : :: : : : : |
| Db 323 ysikskwtvpffkpgvpylykulvlnpdgspasgvp-i-kv-sfsfd-ssgnwitqkrkt | ::: : : : : : : : : :: ::: : : :: : :: ::: ::: ::: :::: ::: :::: ::: |
| 293 | 136 |
| Db 265 feggeamhtl-rgkhile-gypdpklllggslyveasvissdageiensilddipivasp | : ::: : : : : :: : : :: ::: : |

#journal
#title #authors ACCESS IONS

REFERENCE

RESULT ENTRY TITLE ORGANISM DATE

g

à

ð В 윤 ð

Š

윤

셤

ð

셤

ð 유 8

ð

유

유

8

g

ò

원

ð

δ

#authors ACCESSIONS

REFERENCE

TITLE ORGANISM

DATE

RESULT ENTRY CLASSIFICATION SUMMARY

Query Match

셤 ð 염

37

US-08-487-283A-2.rpr

| : ::::: : : : : : :::: : : : | 131 keiavsveimtpenitifreivnpdkgvksggfklpdivsfgtwhvvtrfgstpgktfss 190 | S-FPD | 191 efevkeyvlpsfevsltpakaff-yvddhdltvditarylygkevtgtg-yvvfgvi-t- 246 | 216 YEEVKEYVLZHESVSIEPERNFIGYKNFKNFELITIKARYFYNKVYTEADVYITFGIREDL 275 | 247 tesekksfpaslgrveikdgkgvaclkkeh-itg-tfpkihdlvkgsifvsvsvlteggg 304 | 276 KDDQKEMMQTAMQNTWLINGIAQVTFDSETAVKELSYYSLEDLINKYLYIAVTVIESTGG 335 | 305 emveaekrgiqivtspysilfkrtpkyfkpgmpfdvsvyitnp-dns-pai-gve-v 357 | 336 FSEEAEIPGIKYVLSPYKIALVATPIEIKPGIPYPIKVQVKDSLDQLVGGVPVILNAQTI 395 | 358 evtpd-h-akgvtran-gfakiplntvasatelvitvktkdpgdprqqtgggtmkal 411 | SGVTVLEFNVKTDAPDLPEENQAREGY | 412 pyrtstknfilhvg-vdsn-elkigdpikidlnlgpttipn-hdltymflsrgglvkvg-r 467 | YIDWTDNHKALLVGEHLNIIVTPKSPYIDKITHYNY | 468 fkrggnalvtlsvpvskellpsfrivayyhv-gaadlvadsvwvdikvscmgslkvts 524 | 516 EKFSDASYQSINIPVTQNMVPSSRLLVYYIVTGEQTAELVSDSVWILNIEEKCGNQLQVHL 575 | 525 trpkasyeprrafsltitgdpgakvglvavdkgvyvlnskhrltgtkiwdtiekhdtgct 584 | 576 SPDADAYSPCQTVSLNMATCHDSWVALAAVDSAVYGVQRCAKKPLERVFQFLEKSDLGCG 635 | 585 aggadhmgvfydaglvfetntakgtgirtdpscpvssrrravtisdvitsmasky-hg 643 | NVFHLAGLTFLTHANADDSQENDEPCKETLRPRRTLQKKTEE | lyisdgdvcvqaflvcctemaskkieskqda | 694 VVKKCCYDGACVNNDE-TCEGRAARISLGPRCIKAFTECCVVASQIRANISHKDMQLGRI 752 | 704 eedddddaymrsedivsrsqfpeswmwedtnipecpaqnkhcestsvirnnflkdsittw 763 | 753 HMKTLLPV-SKPE-IRSYFPESWLWE-VHLVV-P-RRKQLQFA-LPDSLTTW 797 | 764 gitaislskthgicvadpfemivlkeffidlklpysavrneglevkailhnysedpiivr 823 | 198 EIGCIGISNT-GICVADTVKAKVFKDVFLEMNIPYSVVRGEQIQIKGTVNYRTSGMQFC 856 | 824 velmengevcssaskkg-kyrqevimdpmstrvvpyviipmklglhsievkasv 876 | | 877 knsgsndgvkrdlrvvaegvlvkketnvllnp-vklgg-eqtshipsgvprnqvpnsdad 934 :::::: | 915 ETWFCKEILVKTLRVVPECVKRESYSCVTLDPRGIYGTISRRKEFPYRIPLDLVPKTEIK 974 | 935 tlisvtagaqtsvlveqaisgdslgslivqpvqcgeqnmiymtlpviat-hyldntkkwe 993 :: | 9/3 RILSVK-GLEVGEILSAVLSQEGINILIHLFKGSAEAELMSV-VEVFIVFHILETGNHMN 1032 | 994 di-gldkrntaikyinigygrqlayrkedgsyaawvsrqsstwltayvvkvfamsstl 1050 |
|--|--|--|--|--|--|--|---|--|--|--|---|--|--|---|--|--|--|--|---------------------------------|--|--|---|--|---|--|--|--|--|---|---|---|
| à | q | δ | QQ | ٥y | QQ | δ | В | ٥y | d d | ٥y | qo | ٥y | QQ | φ | qq | Qy | QQ | δλ | QQ | ð | ΩP | ò | Op | λõ | QQ | Qy | QQ | δ | q | ð | QQ |
| 1180 ihw-gsnkgkaaavestaygilaaiqheegeiaekatnwisqsatfggyfqstqdtvm 1236 | 1233 QHKDSSVPNTGTARMVETTAYALLTSIALKDINYVNPVIKMISEEQRYGGGFYSTQDTIN 1292 | 1237 algaltgfescgsrmkkmdlsfkiraeengvfdkefgitndnafvgkpfkvpvhggltvt 1296 | 1293 AIEGLTEY-SLIVKQLALSMDIDVSYKHKGALHN-YKMTDKN-FLGRPVEVLLMDDLIVS 1349 | 1297 asgtgggiltf-vkkyrekvvikkd-ckgfaleittnldnqvkqrrrqsinpefnvyrfi 1354 | 1350 T-GFGSGLATWHVTTVVHKTSTSEEVC-SFYLKIDTQ-DIEASHYRGYG-NSDYKRIV 1403 | 1355 gcfry-1-rnqep-gmvvmdislptgfeakkkdlddmknlvdnyivqyeirpgrvfly 1409 | SSSHAVMDISLPTGISANEEDLKALVEGVDQLFT | 1410 ldkvnkdekncvgfrlnqvfesnlvlpvtatvfeyyepdfrcsksyhpkmevnpdaschg 1469 : : : :: :: | 1464 LNSIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHRPDKQCTMFY-STSNIKIQKVCEG 1522 | 1470 nicncigrhcvelkgmadedrnad-rngnacraeyvfijgvtkvtktasyjninaalk 1526 | AYAYKVSITSITVENV | 1527 tvikkgmdqainvgarrsfvipmhcgknlnvspqdiylvmgmhnahwr-nsd-rtqyvlt 1584 | AVAEKDSEITFIKKVTC-TNAELVKGRQ | 1585 sdtwfekfplesvc 1598 | 1641 SLTWIETWRRDITC 1654 . | | 101.1 12 151339 #type fragment | ISM | | 151339 151339 | <pre>#authors Lambris, J.D.; Lao, Z.; Pang, J.; Alsenz, J. journal J. Immunol. (1993) 151:1612-6134 ##itla Phical commonst of trust component of the commonst of trust commonst of trust commonst.</pre> | into component of thou comprement. Construing conservation of functional sites. | #CTOSS-felences Mulu:94/003100 #accession 15133 | type | <pre>##residues 1-1620 ##iabei LAM ##ross-references GB:L2433 NID:9431607, CDS_PID:9431608</pre> | DOSKILONION FSUPELLAMILLY Alpha-2-maclogroundi MARY #length 1620 #checksum 2588 | <pre>uery Match 16.1%; Score 1944; DB 6; Length 1620; lest Local Similarity 28.6%; Pred. No. 0.00e+00;</pre> | latches 481; Conservative 421; Mismatches 650; Indels 132; Gaps 89; | 11 pinvkimvknhptqskelasksvvldqannfqamtql·icrgplvddpkqkqyvvlqaqf 70 ::: : : : | 49 AFDATISIKSYPDKKFSYSSGHVHLSSENKFQNSALLTIQPKQLPGGQNPVSTVYLEVVS 108 | 71 pdrllekvvlvsfqsgyifiqtdktiytpastvhyrvfsmtpglepltreifedqevakn 130 |

8 а ð 9 8 원 ð 셤

US-08-487-283A-2.mr

preliminary; translated from GB/EMBL/DDBJ

cross-references MUID:91108039

156095

accession

##molecule_type DNA

Campbell, R.D.; Gagnon, J.; Porter, R.R. Biochem. J. (1981) 199:359-370 Amino acid sequence around the thiol and reactive acyl groups of human complement component C4.

fcross-references MUID:82182029 faccession A17265

##molecule_type_protein #fresidues 757,'X',759-771,980-990 ##label HES

512866

*accession

RESULT

authors fjournal

REFERENCE

#title

classical-complement-pathway C3/C5 convertase (EC 3.4.21.43) 1109 qeassvceqsvnslpgsmakavayl-ekrlp-h---lt-npyavam--tsya-l-anag 1157 1386 IEASHYRGYGNSDYKRIVACASYKPSREESSSGSSHAVMDISLPTGISANEEDLKALVEG 1445 1441 -qkrcvkfyhpqreggtlørlclgdvctcaeescs-mqkkgepdv--q-ridkacgagld 1495 1051 isvqenvlctavkwlilntqqpdgifnefapvihaemtgn--vrgsdndasmtafvliam 1108 1158 klnketllkfasp----q-ldh--wpvpg-gygytleatsyallalvkvkafeeagpi 1207 1327 mdktshed-akesfmltie-vlykns--erda-tm-sildiglltgfivdtddlnglskg 1380 1559 YAYKVSITSITVENVFVKYKATL-LDIYKTGEAVAEKDSEITFIKKVTCTNA-ELVKGRQ 1616 1033 IFHSDPLIEKQKLKKKKKEGMLSIMSYRNADYSYSVWKGGSASTWLTAFALRVLGQVNKY 1092 1268 nknqfhtrtdkvnsidkdltvkas-gngeatlsvvtlyyalpeekdsdcesfdlsvtltk 1326 1501 PDKQCTMFY-STSNI-KIQKVCEGAACKCVEADCGQMQEELDLTISAETRKQTACKPEIA 1558 1496 yvykatvvdskltthtdtytvkidlvi-kpgtdegvegknrdfmglaycrealglmggkt 1554 ymimgksedlhrvedkgllgykyvlgegtwieywpsqqectsrdyrevclgidefingit 1614 Yu, C.Y.

J. Immunol. (1991) 146:1057-1066

The complete exon-intron structure of a human complement component C4A gene. DNA sequences, polymorphism, and linkage to the 21-hydroxylase gene. A17265; A32335; A27600; I58991; I37399; A01262; A01263 #formal name Homo sapiens #common name man 25-Feb-1985 #sequence_revision 23-Aug-1996 #text_change 156095; A29177; B29177; A90845; A19311; A92337; C4b subunit; complement C4a anaphylatoxin complement C4A precursor - human 16-Feb-1997 156095 1615 tfgc 1618 1673 LNGC 1676 13 # journal
#title #authors 1555 **ACCESSIONS** REFERENCE ORGANISM CONTAINS

a

õ

염 8 셤

ð

용

ð

염

à

à

g ð ď

ð

releases the C4a anaphylatoxin from the amino end of the alpha chain and generates C4b, which associates with the 2a fragment of This protein is synthesized as a single-chain precursor and, prior to secretion, is enzymatically cleaved to form a trimer of The activity of C4b is regulated by proteolytic cleavage involving $\mathcal{C}_{\mathbf{Z}}$ Gomez-Escobar, N.; Campbell, R.D. Hum. Mol. Genet. (1994) 3:481-488 Characterisation of the novel gene G11 lying adjacent to the complement G4A gene in the human major histocompatibility Whitehead, A.S.; Goldberger, G.; Woods, D.E.; Markham, A.F.; nonidentical chains (alpha, beta, and gamma) which are linked by complement factor 2 to form the classical-complement-pathway C3 convertase. The C4b, C2a fragment then associates with the 3b 957-1012,'E',1014-1108,'I',1110-1175,'S',1177-1270,'V' 1272-1336 ##label CHA FEBS Lett. (1983) 154:387-390 Amino acid sequence of a polymorphic segment from fragment complement (C4) for analysis of a genetic deficiency of The activation of complement C4 by complement subcomponent C1s Sargent, C.A.; Anderson, M.J.; Hsieh, S.L.; Kendall, E.; The chemical structure of the C4d fragment of the human complement component C4. 1199-1270,'V',1272-1299,'V',1301-1304 ##label CH2 1073-Gly, 1120-Leu, 1121-Ser, 1124-Ile, 1125-His, 1207-Ala, 1210-Arg were also found Use of a cDNA clone for the fourth component of human C4a anaphylatoxin is a vasoactive peptide and a mediator of inflammation. #cross-references EMBL:X77491; NID:g453410; CDS_PID:g453411 Proc. Natl. Acad. Sci. U.S.A. (1983) 80:5387-5391 ##cross-references GB:K00830; NID:g187772; CDS_PID:g187773 Chakravarti, D.N.; Campbell, R.D.; Porter, R.R. Chakravarti, D.N.; Campbell, R.D.; Gagnon, J. ##molecule_type protein ##residues 957-1012,'E',1014-1044 ##latel CAM fragment of complement factor 3 to form the classical-complement-pathway C5 convertase. C4d of human complement component C4 fcross-references MUID:83158189 Mol. Immunol. (1987) 24:1187-1197 translated from GB/EMBL/DDBJ translated from GB/EMBL/DDBJ 1448-1474 ##label RE2 ##molecule_type DNA #fresidues 1-21 ##label RE3 #cross-references MUID:83299979 cross-references MUID:88094444 tcross-references MUID:94282044 in guinea pig. Colten, H.R. ##molecule_type protein ##molecule_type protein disulfide bonds. ##molecule_type mRNA complex A32335 A32335 A27600 158991 137396 158991 137399 #residues ##residues #residues #status #status accession #accession accession #accession ##note #authors # journal | journal #authors | journal #authors **f**authors | journal REFERENCE title REFERENCE ftitle title #title REFERENCE REFERENCE COMMENT COMMENT COMMENT

C4b-binding protein and factor I.

COMMENT

#fcross-references GDB:119732
#map position 6p21.3-6p21.3
#introns 22/2; 88/3; 156/1; 179/3; 209/2; 237/1; 269/2; 304/3; 349/1; 26/2; 34/3; 44/3; 44/3; 44/3; 570/3; 623/3; 666/1; 691/1; 757/1; 794/2; 818/1; 864/3; 934/3; 952/1; 982/1; 1052/1; 1077/2; 1129/3; 1168/3; 1226/1; 1303/3; 1359/3; 1379/3; 1411/1; 1473/2; 1503/3; 1528/3; 1563/1; 1593/1; 1626/1; 1654/1; There are at least two genes coding for C4, C4A and C4B. Each gene Gaps 116; #cross-link thiolester (Cys-Gln) #status experimental/ Residues 1446 or 1449 may be the carboxyl end of the alpha chain. |product C4a anaphylatoxin #status experimental #label product C4d fragment #status experimental #label C4D domain signal sequence fstatus predicted flabel SIGN complement C4 alpha chain #status predicted complement C4 gamma chain #status predicted #cleavage_site Arg-Ala (complement subcomponent Cls)
#status_experimental\ 139 ghlflqtdqpiynpgqrvryrvfaldqkmrpst-dtitvmvenshglrvrkkevympssi 197 198 fq-ddfvipdisepgtwkisarfsdglesnsstqfevkkyvlpnfevkitpgkpyiltvp 256 257 ghldemqldiqaryiygkpv-qgvayvrfgllde--dgkktffrglesqtklvngqshis 313 242 -NFKNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLINGIAQVT 300 314 lskaefqdalekinm-gitdlgglrlyvaaaiieypggemeeaeltswyfvsspfsldls 372 301 FD-SE--TAVKELSYYSLEDIMNKYLY IAVTVIESTGGESERAEIPGIKYVLSPYKIMIN 357 373 ktkrhlvpjapfllqalvremsgspasgipvkvsa-tvsspgsvpevqdiqqntd-gsgq 430 431 vsipiiipqtiselqlsvsagsph-pai--arltvaapp-sg-gpgflsierpds-rppr 484 product complement C4 beta chain #status predicted #superfamily alpha-2-macroglobulin
acute phase; complement classical pathway; glycoprotein; #binding site carbohydrate (Asn) (covalent) #status hydrolase; inflammation; plasma; polymorphism; serine experimental #length 1744 #molecular-weight 192860 #checksum 9431 358 ATPLEIKPGIPYPIKVQVKDSLDQLVGGVPVIINAQTIDVNQETSDLDPSKSVTRVDDGV 466; Conservative 419; Mismatches 620; Indels 147; Length 1744; product C4b #status predicted #label C4B\ fregion C4b-binding protein binding\ Query Match 14.0%; Score 1692; DB 2; I Best Local Similarity 28.2%; Pred. No. 5.65e-280; proteinase; thiolester bond #label BET\ SAM #label product #label product has many alleles. C4A\ 20-675, 757-1446, 1454-1744 CLASSIFICATION 1010-1013 1454-1744 680-1446 757-845 957-1336 680-756 756-757 Matches 20-675 GENETICS #dene KEYWORDS 1 - 19COMMENT FEATURE SUMMARY COMMENT g 셤 අ g ð 셤 ð 음 ð 8 õ

셤

8 6 8 6 8 6

Sy Go

ò

90 yo

gg.

à

Š

q

ò g ð

q ð 셤

US-08-487-283A-2.pr

Jan 28 12:22

| 418 | ASFVIALPSGVTVLEFNVKTDAPDLPEENQAREGYRAIAYSSLSQSYLYIDWTDNHKALL 477 | δ | 1329 | 992 - ' |
|------------|--|-----------------|--------------------------------------|----------------|
| 485 | lravgs-gatfshyywmilsrgqiv-fmnrepkrtltsvsvfv | qq | 1415 | • |
| 478 | | φ | 1372 | _ |
| 541 | fyfvafyy-hgdh-pvanslrvdvqagacegklelsvd-gakgyrngesvklhletds 595 | Db | 1475 | - |
| 538 | raelvsdsvwlniee-kcgnolovhlspdadayspo | Qy | 1414 | - |
| 296 | lalvalgaldtalyaagskshkplnmgkvfeamnsydlgcgpgggdsalqvfqaaglaf- 654 | qū | 1534 | Ψ |
| 597 | DSWVALAAVDSAVY GVQRGAKKPLERVFQFLEKSDLGCGAGGGLANANVFHLAGLTFL 654 | Qy | 1473 | _ |
| 655 | sdgdqwtlerkriscpkekttrkkrnvnfgkaineklgdyasptakrccqdgvtrlpmmr 714 | QQ | 1594 | _ |
| 655 | TNANADDSGENDEPC-KE-ILRPRRTLQKKIEELAAKYKHSVVKKCCYDG-ACVNNDE 709 | Qy | 1531 | _ |
| 715 | | QQ | 1654 | +- |
| 710 | TCEQRAARISLGPRCIKAFTECCVVASQLRA-NISHKDMQLGRLHMKTLLPVSKPE 764 | Qy | 1588 | _ |
| | vreffpenwlwrvetvdrfqiltlwlpdslttweihglslsktkglcvatpvqlrvfref 833 : : : : :::: : : :: | qa | 1711 | Ψ. |
| 765 | IRSYFPESMLMEVHLVPRRKQLQFALPDSLTTWEIQGIGISNT-GICVADTVKAKVFKDV 823 | 0y | 1647 | _ |
| 834 | hlhlrlpmsvrrfeqlelrpvlynyldknltvsvhvspveglclaggglaqqvlvpags 893 : : :: : : : : : : : : FLEMNIPYSVVRGEQ1QLKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKSSKC 883 | RESUL/ ENTRY | RESULT 14 ENTRY | 7 |
| 894 884 | 894 arp-vafsvvptaaaa-slkvvarg-sfefpvg-davskvlqiekegaihreelvye 947 | CON | CONTAINS | |
| 948 | <pre>Inpldhrgrtleipgnsd-pnmip-dgdfnsyv-rvtasdpldtlgse-ga-lspggvas 1002 </pre> | ACCE | DATE ACCESSIONS | |
| 1000 | 1003 llrlprgcgeqtmiylaptlaasryldktegwstlppet-kd-havdliqk-gymriggf 1059 : | | REFERENCE #authore #journal | 99 77 |
| 1060 | 1060 rkadgsyaawl <i>sidsst</i> wltafÿlkvlslaqeqvggspeklqetsnwllsq-qqadgsfq 1118 | | #cross-r #accessi ##mol | × |
| 1119 | | REFI | ##res ##crc ##exp REFERENCE | и о д |
| 1177 | 177 sflgekasagllgahaaaitayaltlt-kapvdllgvahnnlmamaqetgdn-lywgsvt 1234 | - | #authors #journal #title | 8 7 |
| 1235 | 1235 gsqsnavsptpaprnpsdpmpqapalwiettayallhlllhegkaemadqaaawltrggs 1294 | - | #cross-r #accessi | |
| 1295 | 1295 fqqqfrstqdtvialdalsaywiashtteerglnvtlsstgrnqfkshalqlnnrqirgl 1354 | 200 | ##Cro | , , |
| 1355 | l355 eeelqfslgskinvkvggnskgtlkvlrtynvldmknttcqdlqievtvkghveytmean 1414 : :: : : : :: | | #authors | CO . |

| Qy Db | 1329 | TDK-NF-LGRPVEVLLND-D-LIVSTGFGS-GLATVHV-TTV-VH-K-T-STS 1371 edyedyeydelpakdqbdaplqpvtplqlfegrnnrrreapkvveeqesrvhytvciwr 1474 : : :::::::::::::::::::::::::: |
|------------|--|--|
| δy | 1372 | EA-SHYRGY-GNSDY |
| DP Qy | 1475 | ngkvgløgmaiadvtllsgfhalradlekltslædryvshfetegphvllyfdævptsr- 1533 :: : : : : : :: : : : : ESSSG-SSHAVMDISLPTGISANEEDIKALVEGVDQLFTDYQIKDGHVILQINSIPSSDF 1472 |
| qa XX | 1534 | ecvgfeavqevpvglvqpasatlydynperrcsvfygapsksrllaticsaevcqcaeg 1593 |
| 셤 | | kcprqrralerglqdedgyrmkfacyyprveygfqvkvìredsraafrlfetkitqvlhf 1653 : : : : : : |
| δ | 1531 | DCGGMQEELDLTI-SAET-RKQTACK-PEIAYAYKVSITSITVENVFVKYKATLLDIYKT 1587 |
| g & | 1654 | tkdvkaaanqmrnflvrasc-rlrlepgkeylimgldgatydleghpqylldsnswie 1710 ::: : :: :: :: : GEAVAEKDSEI-TFIKKVTCTNAELVKGRQYLIMGKEALQIKVNFSFRYIYPLDSLTWIE 1646 |
| oy Oy | | |
| REA TIT | RESULT 1 ENTRY TITLE CONTAINS | 14 A24558 #type complete complement C4 precursor - mouse classical-complement-pathway C3/C5 convertase (EC 3.4.21.43) C4b subunit; complement C4a anaphylatoxin |
| ORGAL | ORGANISM DATE | #formal name Mus musculus #common name house mouse 31-Mar-1989 #sequence_revision 11-Nov-1994 #text_change 13-Mar-1997 |
| ACC | ACCESSIONS | A2 |
| REE | REFERENCE | A24558 A24558 A24558 A24558 A2558 A2 |
| | <pre># authors # journal # title</pre> | s Septem, U.S.; Noblain, D.S.; Ogdes, K.I. Complete CDNA sequence of the fourth component of murine |
| | #cross- | complement. fcross-references MUID:85298264 |
| | #accession ##molec | cession A2458 ##molecule type mRNA |
| | ##re | ##residues |
| i | ##ex | #fexperimental source strain B10.WR |
| E E | REFERENCE #authors | |
| | # journar #title | |
| | #cross-ref | eren |
| | # E | <pre>##molecule_type mkNA ##residues</pre> |
| REF | ##cr ##ex REFERENCE | #fcross-references GB:M11789 #fexperimental gource strain FM NCE A94013 |
| | #authors | |
| | | |

##cross-references EMBL:X05314; NID:g50241; CDS_PID:g50242 fcross-references MUID:86176748 ##molecule_type DNA #residues #authors title REFERENCE title #title REFERENCE REFERENCE COMMENT COMMENT Levi-Strauss, M.; Tosi, M.; Steinmetz, M.; Klein, J.; Meo, T. Proc. Natl. Acad. Sci. U.S.A. (1985) 82:1746-1750 Multiple duplications of complement C4 gene correlate with Isolation of cDNA clones specifying the fourth component of Taillon-Miller, P.A.; Shreffler, D.C. J. Immunol. (1988) 141:2382-2387 Structural basis for the C40.1/C40.2 serologic allotypes of cDNA clone spanning the alpha-gamma subunit junction in the H-2-controlled testosterone-independent expression of its precursor of the murine fourth complement component (C4). Tosi, M.; Levi-Strauss, M.; Duponchel, C.; Meo, T. Philos. Trans. R. Soc. Lond. (1984) 306:389-394 Sequence heterogeneity of murine complementary DNA clones related to the C4 and C4-S1p isoforms of the fourth mouse complement and its isotype, sex-limited protein.

*cross-references MUID:85038607 Eur. J. Immunol. (1990) 20:1607-1610 C4 from C4-high and C4-low mouse strains have identical Ogata, R.T.; Shreffler, D.C.; Sepich, D.S.; Lilly, S.P. Proc. Natl. Acad. Sci. U.S.A. (1983) 80:5061-5065 1105-1118,'A',1120-1189,'T',1191-1449 ##label LEV Proc. Natl. Acad. Sci. U.S.A. (1984) 81:6822-6826 #fresidues - . 651-719, 'G', 721-738, 'AI', 741-805 #flabel NO2 #fcross-references GB:M12970 sequences in the region corresponding to the ##molecule_type mRNA ##residues 1360-1400,'S',1402-1511 ##label OG2 ##molecule_type DNA ##residues 961-1205,'Q',1207-1290 ##label TAI isotype-specific segment of human C4 murine complement component C4. fcross-references MUID:89009745 faccession A30520 sex-limited isoform, C4-Slp. 1258-1376 ##label TOS 1099-1142 ##label OGA ##experimental_source strain B10.W7R NCE A93753 ##experimental source strain B10.BR Ogata, R.T.; Zepf, N.E. complement component ##experimental source strain FM SNCE A30520 ##cross-references GB:M23186 fcross-references MUID: 90353398 faccession A60227 ##cross-references GB:K00019 ##cross-references GB:X55493 fcross-references MUID:85166208 ##cross-references GB:K02798 ##cross-references GB:K02798 #cross-references MUID:83273751
#accession A01264 ##molecule_type mRNA ##molecule type mRNA ##molecule_type mRNA ##molecule_type DNA A21692 A22039 A29059 A22039 A01264 ##residues ##residues # residues ##residues accession accession accession **f**authors authors authors | journal authors | journal f journal **f**authors | journal | journal | journal ftitle title REFERENCE #title REFERENCE title #title title REFERENCE REFERENCE REFERENCE

K.; Takahashi, M.
Proc. Natl. Acad. Sci. U.S.A. (1986) 83:7883-7887
Identification of the 5'-flanking regulatory region responsible for the difference in transcriptional control Proc. Natl. Acad. Sci. U.S.A. (1984) 81:4908-4911 Genes for murine fourth complement component (C4) and sex-limited protein (S1p) identified by hybridization to Nonaka, M.; Kimura, H.; Yeul, Y.D.; Yokoyama, S.; Nakayama, Hemenway, C.; Kalff, M.; Stavenhagen, J.; Walthall, D.; ##cross-references GB:M14225; NID:g199291; CDS_PID:g554211 preliminary; translated from GB/EMBL/DDBJ between mouse complement C4 and S1p genes #cross-references MUID:87017050 ##residues 1360-1400,'S',1402-1511 ##label OG3 ##cross-references GB:K00019 C4- and S1p-specific cDNA. fcross-references MUID:84272739 ##experimental source strain B10.W7R NCE I59084 Ogata, R.T.; Sepich, D.S. 1-21 ##label RES ##molecule_type mRNA ##molecule type DNA Robins B41195 148274 159084 ##residues *accession ##status #accession #authors authors journal #authors | journal title REFERENCE

Sequence comparison of alleles of the fourth component of complement (C4) and sex-limited protein (S1p). Nucleic Acids Res. (1986) 14:2539-2554 f journal

591-603, 'M', 605-1323,'N', 1325-1452,'V', 1454-1585,'Q', 1587-1738 ##label RE2 preliminary; translated from GB/EMBL/DDBJ ##molecule_type mRNA 148274 ##residues #status faccession

Nonaka, M.; Nakayama, K.; Yeul, Y.D.; Shimizu, A.; Takahashi 154567

#title Molecular cloning and characterization of complementary and genomic DNA clones for mouse C4 and SLP. #cross-references MUID:86031969 Immunol. Rev. (1985) 87:81-99 f journal

preliminary; translated from GB/EMBL/DDBJ #molecule_type DNA 154567 #status *accession

##cross-references GB:M12968; NID:g199267; CDS_PID:g199270 preliminary; translated from GB/EMBL/DDBJ 1-128 ##label RE3 169023 #accession ##status

This protein is synthesized as a single-chain precursor and, prior to secretion, is enzymatically cleaved to form a trimer of nonidentical chains (alpha, beta, and gamma) which are linked by disulfide bonds. #cross-references GB:M12969; NID:g199268; CDS_PID:g387439 1724-1738 ##label RE4 ##residues

releases the C4a anaphylatoxin from the amino end of the alpha chain and generates C4b, which associates with the 2a fragment of complement factor 2 to form the classical-complement-pathway C3 convertase. The C4b,C2a fragment then associates with the 3b The activation of complement C4 by complement subcomponent C1s

##experimental_source strain B10.W7R

COMMENT

US-08-487-283A-2.mr

1002 1058 1117 1122 1175 1175 1292 1352 598 655 714 711 773 833 893 1233 1330 g Š g ð 9 ð අ Š 염 à 셤 δ 셤 à a ð ď ð 셤 ð 쉱 S g ð g ð g õ g ð 셤 C4a anaphylatoxin #status predicted #label C4A\ activity of C4b is regulated by proteolytic cleavage involving Gaps 111; #{domain signal sequence #status predicted #label SIG\
product complement C4 beta chain #status predicted
#label BET\ 22/2; 86/3; 977/3; 1047/3; 1073/3; 1125/3; 1164/3; 1221/3 the list of introns is incomplete complement C4b #status predicted #label C4B\ 483 gdtfilnlqpvgjpaptfshyyymiisrgqimamg-rep-rktv-tsvsvlvdhqlapsf 539 |: : : : | : : | |: | |:||:|| |: | || complement C4 alpha chain #status predicted | :|::|| |:| |:| | |: ||::|: ::|: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || :: || snsdeiqldigaryiygkpv-qgvaytrfalmde--qgkrtflrgletqaklvegrthis 311 429 sisfpipptvtelrllvsagsl-ypai--arltvqa-p-psrgtgflsie-pldprspsv 482 complement C4 gamma chain #status predicted 195 ifqdaftipdisepgtwkisarfsdglesnrsthfevkkyvlpnfevkitpwkpyilmvp 254 241 KNFKNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLINGIAQVT 300 312 iskdqfqaaldkinigvrdleglrlyaataviespggemeeaeltswrfvssafsldlsr 371 372 tkrhlvpgahfllqalvqemsgseasnvpvkvsa-tlvsgsdsqvldiqqstngigq-v- 428 419 SFVIMIASGVTVIEFNVKTDAPDIAEENQARECYRAIAYSSLSQSYLYIDWTDNHKALLV 478 137 ghifvqtdqpiynpgqrvryrvfaldqkmrpstdfltitvenshglrv-lkkei-ftsts 194 cleavage_site Arg-Asn (complement subcomponent Cls) acute phase; complement classical pathway; glycoprotein; hydrolase; inflammation; plasma; polymorphism; serine !binding_site carbohydrate (Asn) (covalent) #status product C4d fragment #status predicted #label C4D\ #cross-link thiolester (Cys-Gln) #status predicted
#length 1738 #molecular-weight 192870 #checksum 4149 ö classical-complement-pathway C5 convertase. C4a anaphylatoxin is a vasoactive peptide and a mediator 449; Conservative 407; Mismatches 628; Indels 143; DB 2; Length 1738; region C4b-binding protein binding/ fragment of complement factor 3 to form the Query Match 13.3%; Score 1605; DB 2; I Best Local Similarity 27.6%; Pred. No. 1.46e-263; fsuperfamily alpha-2-macroglobulin C4b-binding protein and factor I. proteinase; thiolester bond #status predicted/ ALP #label GAM\ predicted\ product product product product | label inflammation. 20-673, 754-1440, The The 224, 743, 1387 CLASSIFICATION 1448-1738 1006-1009 1448-1738 #introns 678-1440 953-1332 678-753 154-843 753-754 Matches 20-673 255 note GENETICS KEYWORDS

SUMMARY

용 8 셤 5 셤 ð 용 ð 원 8 용 8 셤 8

1116 1000 1411 1057 1232 1228 1329 1001 1061 1121 1174 qdkvvlrptaprsptepvpqapalwiettayall-hlllregkgkmadkaaswlthqgsf 1291 1372 1412 dyedyydmpaaddpsvplqpvtplqlfegrrsrrreapkvaeeqesrvqytvciwrngk 1471 EVCSFY-LKI-DTQDI--EA-SHYRGY-GN-SDYKR-I--VA----C-ASYKPS-REESS 1416 1351 655 713 710 772 765 832 824 892 884 943 540 yfvayfyhqg-h-p--vanslliniqsrdcegklqlkvd-gakeyrnadmmklriqtdsk 594 :: |: |: | : | : | |: | 597 654 SHVALAAVDSAVYGVQRGAKKPLE--RVFQFLEKSDLGCGAGGGLNNANVFHLAGLTFLT ceqraarvpqqa-crepflscckfaedlrrnqtrsqahlarnnhnmlqeedlideddilv LIENTIPAQ-STFTLAISAYALSLGDKTHPQFRSIV-SALKREALVKGNPPIYR---F-W hgafrstqdtvvtldal8aywiashtteekalkvtl8smgrnglkthglhlnnhqvkgle 539 RLLVYYIVTGEQTAELVSDSVWINIEEK-CGNQLQVHLSPDADAYSPGQTVSINMATGMD 595 alvalgavdtalyavggrshkpldmskvfevinsynvgcgpggddalqvfgdaglafsd gdrltgtre-dlscpkekksrgkrnvnfgkavseklggysspdakrccgdgmtklpmkrt Ihlrlpisirrfeqfelrpvlynylnddvavsvhvtpveglclagggmmaggvtvpagsa rp-vafsvvptaaanv-plkvvargv-fdl----g-davskilqiekegaihreelv-yn 944 ldp--1-nnlgrtleipgssdpnivpdgdfsslvrvtasepletmgsegalspggvasll 944 LDPRGIYGTISRRKEFPYRIPLDLVPKTEIKRILSVKGLLVGEIL-S-AVLSQEGINILT 1001 rlpqgcaeqtmiylaptltasnyldrteqwsklspet-kd-havdliqk-gymriqqfrk HIPKGSAEAELMSVVPVFYVFHYLETGNHWNIFHSDPLIEKQKIKKKLKEGMLSIMSYRN lgqkasagllgahaaaitayaltlt-kasedlrnvahnslmamaeetgeh-lywglvlgs 1281 GGGFYSTQDTINAIEGLTEY---SLLV--KQLRLSMD-ID-VSYKHKGA-LHNY--K-MT 656 NANADDSQENDEPC-KEIL-RPRRTL--QKKIEEIAAKYKHSVVKKCCYDGAC-VNNDET rtsfpenwlwrvepvdssklitvwlpdsmttweihgvslskskglcvakptrvrvfrkfh cpvihramqggl-vgs-detvaltafvvialhhgldvfqdddakqlknrveasitkansf SQYQPIKLQGTLPVEARENSLYLTAFTVIGIRKA---F--DIC-PL-VKIDTALIKADNF eelkfslgstisvkvegnskgtlkilrtynvldmknttcqdlqievkvtgaveyawdane 885 RQKVEGSSSHLVTFTVLPLEIGLHNINFSLETWFGKEILVKTLRVVPEG-VKRESYSGVT --_ = = .. = :: -----:: :: :: - .. :: 1373 à

| 1116 | |
|-----------------|-----------|
| dpcpv | |
| 뜌 | _ |
| 25 | = |
| ъ. | ••• |
| 5 | = |
| ੰਦੇ | _ |
| llaq- | <u></u> |
| 3 | _ |
| ptas | :: |
| ĭ | |
| -8 | |
| <u>,</u> | |
| ű | •• |
| 5 | |
| ıqeq, | _ |
| - | |
| 18 | <u></u> |
| -::1 | •• |
| = | <u></u> |
| 2 | = |
| ď | _ |
| ÷ | = |
| 3 | _ |
| Bt | = |
| स | ••• |
| ĕ | |
| ₹ | |
| 76 | _ |
| Ę | <u>::</u> |
| S | |
| 1058 sfgawlhrds | |
| qq | |

| | =# |
|----------|---|
| | 1124 |
| | |
| • | 1065 SYSVWKGGSASTWLTAFALRVLGQVNKYVEQNQNSICNSLLWLVENYQLDNGSFKENSQY |
| _ | Ĕ |
| - | ĕ |
| • | ą |
| _ | ĕ |
| | VENYO |
| • | 볼 |
| = | 불 |
| | 3 |
| : | S |
| ٠ | 음 |
| • | S |
| • | ž |
| | ជ |
| _ | 2 |
| | ž |
| | 8 |
| • | WLTAFALRVLG |
| : | ≳ |
| _ | ᆿ |
| <u>-</u> | € |
| - | < |
| _ | S |
| - | |
| _ | 22 |
| ٠ | Α. |
| | 8 |
| | ខ |
| _ | WKGGSAS |
| | S |
| : | Ξ. |
| _ | ŝ |
| | 65 |
| | õ |
| | 1 |
| | |

ð

1652 TTCSS 1656

δ

g

Š

요

Š

요

Š g

¹⁵⁹¹ rslerrvedkdgyrmrfacyyhqveygftvkvlredgraafrlfeskitqvlhfrt-dtm 1649 셤

ð 셤

ð

¹⁷⁰⁷ qucks 1711 g

Search completed: Wed Jan 28 12:15:48 1998 Job time : 99 secs.

US-08-487-283A-2.pp

a G 8 ď 셤 염 g 엄 쇰 q 쇰 ð ð ð ð õ ð 염 g g 염 요 g à Š Š ð Š ð Š ð 22/2; 86/3; 154/1; 177/3; 207/2; 235/1; 267/2; 302/3; 347/1; 385/3; 444/3; 506/3; 568/3; 621/3; 664/1; 689/1; 754/1; 789/2; 813/1; 859/3; 927/3; 945/1; 975/1; 1045/1; 1070/2; 1122/3; 1161/3; 1219/1; 1296/3; 1352/3; 1372/3; 1404/1; 1464/2; 1494/3; 1519/3; 1554/1; 1584/1; 1617/1; 1645/1; 1417 SG-SSHAVMDISLPTGISANEEDLKALVEGVDQLFTDYQIKDGHVILQLNSIPSSDFLCV 1475 1531 gfgasqevvvglvqpssavlydyyspdhkcsvfyaaptksqllatlcsgdvcqcaegkcp 1590 1591 rllrslerrvedkdgyrmrfacyyprveygftvkvlredgraafrlfeskitqvlhfrkd 1650 1651 tmasigqtrnf1srasc-r1rlepnkeylimgmdgetsdnkgdpqyl--1dsntwieemp 1707 1472 lglsgmaiaditllsgfhalradlekltslsdryvshfetdgphvllyfdsvpttr-ecv 1530 that sex-limited ŏ 27-Oct-1995 #sequence_revision 03-Nov-1995 #text_change Hemenway, C.; Kalff, M.; Stavenhagen, J.; Walthall, D.; component formal name Mus musculus #common name house mouse #superfamily alpha-2-macroglobulin
#length 1735 #molecular-weight 192820 #checksum ##residues 634-641, L', 643-828 ##label RES ##cross-references EMBL:X06455, NID:954102; CDS_PID:9899336 y Match 13.0%; Score 1565; DB 6; Length 1735; Local Similarity 27.2%; Pred. No. 5.00e-256; fjournal Nucleic Acids Res. (1986) 14:2539-2554
fittle Sequence comparison of alleles of the fourth co-complement (C4) and sex-limited protein (S1p) fcross-references NUID:86176748
faccession 148770 preliminary; translated from GB/EMBL/DDBJ Ogata, R.T.; Zepf, N.E. J. Immunol. (1991) 147:2756-2763 The murine Slp gene. Additional evidence preliminary; translation not shown sex-limited protein Slp(w7) - mouse protein has no biologic function #type complete 1-1735 ##label OGA ##cross-references EMBL:M64933 13-Mar-1997 S54784; I48770 Robins, D. ##molecule_type mRNA ##molecule_type DNA 1689/2 **S54784** 554784 148274 1708 sequecks 1714 1650 RDTTCSS 1656 ##residues #status CLASSIFICATION ##status *accession Query Match fjournal ftitle fauthors | #authors #introns ACCESS IONS TITLE ORGANISM REFERENCE REFERENCE GENETICS RESULT ENTRY 용 ð g ð 쉽 원 q ð δ δ

420; Mismatches 621; Indels 142; Gaps 111;

442; Conservative

Best Loc Matches

944 --1-nnlgqmleipgssdpnivpegdfstfvkvtasepletlgsegalspggvasl1rlp 1000 rscaeqtmiylaptltasnyldrtkqwsklspet-kd-havhliqk-ghvriqqfrkkdg 1057 :: || :: : | : : ||: :|: :|: | 1004 137 ghifvqtdqpiynpgqpvryrvfaldqkmrpstdfltitvenshglrv-lkkei-ftats 194 | :|::|| |:| |:| |:| |::| |: |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: || |: | snsdeigldigaryiygkpv-ggvaytrfalmde--ggkrtflrgletgaklvegrthis 311 | :::: | | | | | | | | | | | :: | | | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: | ::: 371 358 428 | | | | | : : | :: | | :: | | :: | | :: | TPLFLEAPGIPYPIKVQVKDSLDQUSCVPVILNAQTIDVNQETSDLDPSKSVTRVDDGVA 418 892 vafsvvptas-tnvplkl-vak-g-tlds--sdsv-skilqiekegaihreeiv-ynldp 943 946 1005 KGSAEAELMSVVPVFYVFHYLETGNHWNIFHSDPLIEKQKLKKKKKEGMLSIMSYRNADY 1064 594 597 654 655 gdrltqtre-discpkekksrqkrnvnflkalseklgqysspdakrccqdgmtklpmkrt 713 656 NANADDSQENDEPC-KEIL-RPRRTLQ--KKIEEIAAKYKHSVVKKCCYDGAC-VNNDET 710 827 887 833 rlpisvrrfeqlelrpvlynylnddknvsvhvtpveglcmagggmmagwvivpagsalp- 891 RGIYGTISRRKEFPYRIPLDLYPKTEIKRILSVKGLLVGEIL-S-AVLSQEGINILTHLP tkrhlvpgahfllgalvreisgseasnvivkvsa-tlvsgsdsgvlnvggstnrigg-vgdtfilnlqavgipaptfshyyymiisrgqimam-srearrtvt--svsvlvdhqlapsf 479 GEHLNIIVTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPSS 595 alvalgavdtalyavggwshkpldmskvfevidsynlgcgpggddapqvfqdaglafsd fpdnwlwrvepvdssklltvrlpdsmttweihgvslskskglcvakptrvrvfrkfhlhl 828 NIPYSVVRGEQIQIKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKSSKCVRQK 888 VEGSSSHLVTFTVLPLEIGLHNINFSLETWFGKEILVKTLRVVPEG-VKRESYSGVTLDP <u>..</u> ----255 241 312 301 372 359 483 655 773 1001 a ð

US-08-487-283A-2.rsp

| | (MT) |
|--|-------------|
| | :- <u> </u> |
| | |
| | |
| | |
| | |
| | |
| | <u> </u> |
| | |

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 University of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

protein - protein database search, using Smith-Waterman algorithm MPsrch_pp

Wed Jan 28 12:16:06 1998; MasPar time 37.99 Seconds 935.704 Million cell updates/sec Run on:

Tabular output not generated.

>US-08-487-283A-2 (1-1676) from US08487283A.pep 12048

Perfect Score: Description:

1 MGLLGILCFLIFLGKTWGQE..........CQAFLANLDEFAEDIFLNGC 1676 Sequence:

PAM 150 Gap 11 Scoring table:

59021 seqs, 21210388 residues

Searched:

Listing first 45 summaries Post-processing: Minimum Match 0%

1:part1 2:part2 3:part3 4:part4 5:part5 6:part6 7:part7 8:part8 9:part9 10:part10 11:part11 swiss-prot34

Database:

Mean 58.177; Variance 98.799; scale 0.589 Statistics:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| | Pred. No. | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 0.00e+00 | 5.45e-148 |
|-----|--------------------------|------------|------------|--------------|--------------|--------------|--------------|--------------|--------------|--------------|-------------|-------------|-------------|--------------|
| | | PRECURS | PRECURS | PRECURS | PRECURS | PRECURS | PRECURS | PRECURS | PRECURS | (CONTAI | 33-1 (CONT | PRECURS | C4 PRECURS | ALPHA C |
| | ~ | 5 | S | \mathbb{S} | _ | | 2 | \mathbb{S} |
| | Description | COMPLEMENT | COMPLEMENT | COMP LEMENT | COMPLEMENT | COMPLEMENT | COMP LEMENT | COMP LEMENT | COMP LEMENT | COMP LEMENT | COMP LEMENT | COMP LEMENT | COMP LEMENT | COMPLEMENT |
| | 10 | CO5 HUMAN | CO5_MOUSE | CO3 NAJNA | CO3_HUMAN | CO3_LAMJA | CO3_MOUSE | CO3_CAVPO | CO3_RAT | CO3_EPTBU | CO3 ONCMY | CO4 HUMAN | CO4_MOUSE | CO3_RABIT |
| | 98 | 7 | 7 | 7 | ~ | 2 | 7 | 7 | 7 | 7 | 7 | ~ | 7 | ~ |
| | Query Match Length DB | 1676 | 1680 | 1651 | 1663 | 1673 | 1663 | 1666 | 1663 | 1620 | 1640 | 1741 | 1738 | 126 |
| مين | Query Match | 100.0 | 81.3 | 19.6 | 19.6 | 19.2 | 19.1 | 19.1 | 18.7 | 17.5 | 16.6 | 14.3 | 13.3 | 9.9 |
| | Score | 12048 | 9789 | 2367 | 2361 | 2309 | 2296 | 2302 | 2255 | 2106 | 2002 | 1723 | 1605 | 196 |
| | Result No. | | 7 | 3 | 4 | 2 | 9 | 7 | œ | 6 | 10 | 11 | 12 | 13 |

| 14 | 670 | 9.0 | 1477 | ٦. | Ali3 RAT | ALPHA-1-INHIBITOR III | 2.99e-118 |
|----|-----|---------|------|----|------------|-----------------------|-------------|
| 15 | 199 | 5.5 | 1476 | - | AZMG_MOUSE | MURINOGLOBULIN 1 PREC | 3.83e - 116 |
| 16 | 583 | 4.8 | 1474 | | A2MG HUMAN | ALPHA-2-MACROGLOBULIN | 8.97e-98 |
| 11 | 581 | 4.8 | 1482 | σ | PZP HUMAN | PREGNANCY ZONE PROTEI | 1.53e - 97 |
| 18 | 208 | 4.2 | 323 | 7 | CO3 XENLA | COMPLEMENT C3 (FRAGME | 8.31e - 81 |
| 19 | 207 | 4.2 | 1472 | - | A2MG RAT | ALPHA-2-MACROGLOBULIN | 1.40e-80 |
| 50 | 205 | 4.2 | 1473 | _ | OVOS CHICK | OVOSTATIN PRECURSOR (| 1.92e - 79 |
| 21 | 493 | 4.1 | 1451 | - | A2MH MOUSE | MURINOGLOBULIN 2 PREC | 2.11e-77 |
| 22 | 399 | 3,3 | 74 | 7 | CO5A PIG | COMPLEMENT C5A ANAPHY | 2.09e-56 |
| 23 | 382 | 3.2 | 74 | 7 | CO5A BOVIN | COMPLEMENT C5A ANAPHY | 2.44e-53 |
| 24 | 351 | 5.9 | 16 | 7 | COSA_RAT | COMPLEMENT C5A ANAPHY | 5.68e-46 |
| 52 | 171 | 1.4 | 16 | 7 | CO4A RAT | COMPLEMENT C4A ANAPHY | 3.12e-10 |
| 56 | 170 | 1.4 | 11 | 7 | CO4A BOVIN | COMPLEMENT C4A ANAPHY | 4.64e-10 |
| 27 | 139 | 1.2 | 77 | 7 | CO3A_PIG | COMPLEMENT C3A ANAPHY | 5.44e-05 |
| 88 | 121 | 1.0 | 798 | 10 | YA73_SCHPO | HYPOTHETICAL 92.1 KD | 2.39e-02 |
| 53 | 115 | 1.0 | 1292 | œ | RPOC MYCGE | DNA-DIRECTED RNA POLY | 1.57e-01 |
| 30 | 105 | 6.0 | 257 | Ξ | YGBI HAEIN | HYPOTHETICAL TRANSCRI | 3.00e+00 |
| 31 | 106 | 0.9 | 270 | 11 | YD1J SCHPO | HYPOTHETICAL 31.5 KD | 2.26e+00 |
| 32 | 105 | 6.0 | 328 | | ASCD_YERPS | CDP-6-DEOXY-DELTA-3,4 | 3.00e+00 |
| 33 | 110 | 0.9 | 341 | 11 | YEJK HAEIN | HYPOTHETICAL PROTEIN | 7.10e-01 |
| 34 | 108 | 0.9 | 461 | - | AD4B BOVIN | STEROID HORMONE RECEP | 1.27e+00 |
| 35 | 106 | 0.0 | 462 | - | AD4B MOUSE | STEROID HORMONE RECEP | 2.26e+00 |
| 36 | 112 | 6.0 | 472 | œ | PSBC_SYNY3 | PHOTOSYSTEM II 44 KD | 3.91e-01 |
| 37 | 108 | 6.0 | 589 | S | KPYK LACDE | PYRUVATE KINASE (EC 2 | 1.27e+00 |
| 38 | 108 | 6.0 | 612 | 10 | UNC6 CAEEL | UNC-6 PROTEIN PRECURS | 1.27e+00 |
| 39 | 108 | 6. | 943 | 'n | IROA NEIME | IRON-REGULATED OUTER | 1.27e+00 |
| 40 | 113 | 6.0 | 1024 | 10 | UBA1 YEAST | UBIQUITIN-ACTIVATING | 2.90e-01 |
| 41 | 106 | ō: 0 | 1030 | 10 | VPP1 CAEEL | PUTATIVE CLATHRIN-COA | 2.26e+00 |
| 42 | 112 | 6.0 | 1517 | 4 | GLTB ECOLI | GLUTAMATE SYNTHASE (N | 3.91e-01 |
| 43 | 107 | 6.0 | 1736 | Ξ | ZO1 HUMAN | TIGHT JUNCTION PROTEI | 1.70e+00 |
| 44 | 108 | 6.0 | 1805 | 10 | Y218 MYCGE | HYPOTHETICAL PROTEIN | 1.27e+00 |
| 45 | 106 | 6:0 | 5147 | m | FAT DROME | CADHERIN-RELATED TUMO | 2.26e+00 |
| | | | | | ļ | | |

ALI GNMENTS

| | | | | | | | | | | | | | | R.A.; | | | | | | | | | | | | |
|----------|---------------|---------|--------------------------------|-------------|---|--|-----|-----------------------|--|---------------------|----|--------------------|--------------------|---------------------------------------|--------------------------------|-----|--------------------------------|--------------------|---|------------|-----------------------------------|-----|-------------------------------|--------------------|--|---|
| | | | | | | LATOXIN). | | | EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; | | | | | FLEISCHER D.T., HUNT A., WETSEL R.A.; | | | | | WETSEL R.A., LEMONS R.S., LEBEAU M.M., BARNUM S.R., NOACK D., | | | | | | LUNDWALL A.B., WETSEL R.A., KRISTENSEN T., WHITEHEAD A.S., | •• |
| | Ā | | | ධ | ATE) | APHY | | | TET | | | | | : | | | | | RNUM | | | | | | H | В. F |
| | PRT; 1676 AA. | | | UPDAT | ON UPD | CSA AN | | | BRATA; | | | | | CHER D | | | | | М., BA | | | | | | SEN T. | , TACK |
| | PRT; | | | UENCE | TATIO | INS: (| | | VERTE | | | | | | | | | | CAU M.P | | <u>.</u> | | | | (ISTEN | I H.R., |
| | 83; | | CREATED) | LAST SEC | LAST AN | JR (CONTA | | | HORDATA; | | | | | ND J.C., | 68 (1991). | | FROM N.A. | | .S., LEBE | | 1482 (1988 | | ROM N.A. | | R.A., KF | ., COLTER |
| | STANDARD; | | 01, | 24, | 33, | CURS | | ₹ | C | ۲. | | | | /ILM | 62-3 | | 919 | | NS R | | 174-1 | |)2 FI | | FSEL | R.C |
| | ST | | (REL. | (REL. | (REL. | 5 PRE | | (HOM) | ETA207 | IMATE | | M N.A | 79575 | ., HA | 146:30 | | 412-10 | 09511 | LEMO | | 27:14 | | 412-9(| 30937. | . WE | OCDEN |
| KESULT 1 | CO5_HUMAN | P01031; | 21-JUL-1986 (REL. 01, CREATED) | 01-DEC-1992 | 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) | COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN). | c5. | HOMO SAPIENS (HUMAN). | EUKARYOTA; M. | EUTHERIA; PRIMATES. | El | SEQUENCE FROM N.A. | MEDLINE; 91079575. | HAVILAND D.L., HAVILAND J.C., | J. IMMUNOL. 146:362-368(1991). | [2] | SEQUENCE OF 412-1676 FROM N.A. | MEDLINE; 88209511. | WETSEL R.A., | TACK B.F.; | BIOCHEMISTRY 27:1474-1482 (1988). | [3] | SEQUENCE OF 412-902 FROM N.A. | MEDLINE; 85130937. | LUNDWALL A.B | WOODS D.E., OGDEN R.C., COLTEN H.R., TACK B.F.; |
| 200 | Ω | ပ္ | Ħ | 5 | Ţ. | 띩 | A | S | ဗ | ဗ | ≅ | ₽ | × | \$ | .7 | 3 | ₽ | ⋨ | \$ | ≴ | ₽ | ⋧ | ۵ | ⋨ | ≴ | ≴ |

J. BIOL. CHEM. 260:2108-2112(1985).

US-08-487-283A-2.rsp

m

F -> S. MW; 9D5C6E59 CRC32; POTENTIAL. POTENTIAL POTENTIAL 1676 AA; 188331 731 732 741 911 1115 1630 518 699 711 741 911 1115 1630 518 DISULFID DISULFID CARBOHYD CARBOHYD CARBOHYD CARBOHYD SEQUENCE VARIANT FFFFF

Gaps ö Length 1676; Indels ö 5; Pred. No. 0.00e+00 Score 12048; DB 0; Mismatches 100.0%; Similarity 100.0%; Matches 1676; Conservative Query Match Best Local

ö

180 61 DKKFSYSSGHVHLSSENKFQNSALLTIQPRQLPGCQNPVSTVYLEVVSKHFSKSKRMPIT 120 121 ydngflfihtdkpvytpdqsvkvrvyslnddlkpakretvltfidpegsevdmveeidhi 셤 ŝ

240 181 giisfpdfkipsnprygmwtikakykedfsttgtayfevkeyvlphfsvsiepeynfigy 용

ð

180

240 300 181 GIISFPDFKIPSNPRYGMWTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFIGY knfknfeitikaryfynkvvteadvyitfgiredlkddqkemmqtamqntmlingiaqvt 241 g ð

300 241 KNFKNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLINGIAQVT ð

301 fdsetavkelsyysledlnnkylyiavtviestggfseeaeipgikyvlspyklnlvatp 360 301 FDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKINLVATP 360 염 ð 420 420 셤 õ

480 480 vlnlpsgvtvlefnvktdapdlpeengaregyraiaysslsgsylyid*tdnhkallvge 421 g ð

hlniivtpkspyidkithynylilskgkiihfgtrekfsdasygsinipvtgnmvpssrl 481 481 g à

9 9 541 lvyyivtgeqtaelvsdsvwlnieekcgnqlqvhlspdadayspgqtvslnmatgmdswv g δ

99 9 alaavdsavygvqrgakkplervfqfleksdlgcgaggglnnanvfhlagltfltnanad 601 ALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGLNNANVFHLAGLTFLTNANAD 601 a à 720 720 dsqendepckeilrprrtlqkkieeiaakykhsvvkkccydgacvnndetceqraarisl DSQENDEPCKEILRPRRTLQKKIEEIAAKYKHSVVKKCCYDGACVNNDETCEQRAARISL 661 661 g õ 721 721

원 ò

| • | | |
|---------------------|---------|-------------|
| | ø | |
| | 0 | |
| | | |
| 90 | | |
| | ٠. | |
| • | | |
| | 43 | |
| | • | |
| | ••• | • |
| | ٠. | |
| ••• | | |
| | | |
| | | |
| 00 | ••• | |
| | ٠. | |
| ٠., | ٠. | |
| • | | |
| | | |
| | • | |
| ٠., | ٠., | • |
| ٠. | | |
| ••• | ж. | |
| | | |
| | • | |
| | • | |
| | • | • |
| ٠. | ٠. | |
| | • | |
| | • | |
| 00 | • | |
| | ٠. | |
| • | ٠. | |
| • | | |
| | 0 | |
| | | • • • |
| | - | |
| | ď | |
| | -2 | |
| | ٧. | |
| | | |
| | | |
| | | |
| × | ٠. | |
| | | |
| | | |
| œ | ٠. | |
| | ٠. | 900 |
| | - | 200 |
| | æ | 100 |
| | | 200 |
| œ | c | |
| - | ٠, | |
| ж. | | |
| | | |
| • | ٠, | |
| ٠. | ٠. | |
| - | | |
| | | |
| | | |
| | 2 | - |
| ÷ | Ť | ж |
| × | Ĭ | |
| Š | Ĭ | |
| ž | į | |
| 9 | į | |
| | Ĭ | |
| 200 | Ĭ | |
| 90 | Ĭ | , |
| A 00 . | Ĭ | |
| 24.00 | į | • |
| 10 AC AC | Ĭ | |
| 10 AC | Ĭ | |
| 110 00 40 | 1577 | • |
| 110 00 40 | Ť | |
| 110 00 40 | Ĭ | |
| 110 A 00 A 00 A 011 | To To | |
| 110 00 11 | | |
| 110 00 1 | | |
| 110 00 1 | |) |
| 110 00 11 | | |
| 37 06 611 | | |
| 37 06 611 | | · · |
| 37 00 CII | |) |
| 37 0V CII | |))) |
| 110 00 41 | | ; ; |
| 110 00 41 | | · · |
| 110 00 41 | Tara Ca | ; ; |
| 37 OV G11 | |) |
| 37 0V CII | | · · |
| 110 00 41 | |) |
| 110 00 41 | | 1 |
| 37 00 CT1 | |) |
| 37 00 011 | | ; ; ; |
| 37 00 011 | | ; ; |
| 37 DV C11 | | ; ; ; |
| 37 0V GI1 | | ; ; ; |
| 37 DV G11 | | ; ; ; |
| 37 00 011 | | · · · |
| 37 0V 011 | | ; ; ; |
| 37 04 411 | | ; ; ; |
| 37 OV C11 | | · · |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| | | |
| 37 00 G11 | | |

| | | | | | | | | | , | | | | | |
|--|--|------------|---|--|--|--|--|---|--|---|--|--|--|---|
| 840 | 006 | 096 096 | 1020 | 1080 | 1140 | 1200 | 1260 1260 | 1320 1320 | 1380 1380 | 1440 1440 | 1500 | 1560 1560 | 1620 1620 | |
| prrkqlqfalpdslttweiggigisntgicvadtvkakvfkdvflemnipysvvrgeqig | lkgtvynyrtsgmqfcvkmsavegictsespvidhqgtksskcvrqkvegssshlvtftv | | yripidlypkteikrilsvkgllygeilsavlsqeqinilthlygsaeaelmsvypvfy | vfhyletgnhwnifhsdpliekqklkkklkegmlsimsyrnadysysvwkggsastwlta | falrvlggvnkyvegngnsicnsllwlvenyqldngsfkensgygpiklggtlpvearen | slyltaftvigirkafdicplvkidtalikadnfllentlpagstftlaisayalslgdk | thpqfrsivsalkrealvkgnppiyrfwkdnlqhkdssvpntgtarmvettayalltaln | kdinyvnpvikw seegrygggfystddtinaieglteys vkg rlsmdidvsykhk | gaihnykmtdknflgrpvevllnddlivstgfgsglatvhvttvvhktstseevcsfylk | idtgdieashyrgygnsdykrivacasykpsreessgsshavmdis1ptgisaneedlk | alvegvdqlftdyqikdghvilqlnsipssdflcvrfrifelfevgflspatftvyeyhr | pdkqctmfystsnikiqkvcegaackcveadcgqmqeeldltisaetrkqtackpeiaya | ykvsitsitvenvfvkykatlldiyktgeavaekdseitfikkvtctnaelvkgrgylim | gkealqikynfafryiypldsltwieywprdttcsscqaflanldefaediflngc 1676 |
| 781 | 841 | 901 | 961 | 1021 | 1081 | 1141 | 1201 | 1261 | 1321 | 1381 1381 | 1441 | 1501 | 1561 1561 | 1621 |
| 95 Q | g & | go Oy | do y | g S | g y | 40 y | දු දි | g y | g & | g & | g & | g y | g & | g & |

PRT; 1680 AA

STANDARD;

CO5 MOUSE

RESULT ID CO

US-08-487-283A-2.rsp Jan 28 12:23

S

P06684;

O

```
WETSEL R.A., OGATA R.T., TACK B.F.;
BIOCHEMISTRY 26:737-743(1987).
-!- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE SPONTANEOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9, INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIETT BINDING SITE FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYTIC
                                                                                                                                                                                                                                                                                                                                 SUBUNIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC RESIDUES, FORMING TWO CHAINS, BETA 4 ALPHA, LINKED BY A DISULFIDE BOND. C5 CONVERTASE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN, RELEASING C5A ANAPHYLATOXIN 4 GENERATING C5B (BETA CHAIN + ALPHA'
                                                                                                                                                                                                                                                                                                                                                                                            CHAIN).

SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN.
FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C5,
TOTAL TABLEMANATORY PROCESS. IT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        PIR; A27538; A27538.
PIR; A35530; A35530.
HSSP; POLOZ2; IC5A.
HSSP; POLOZ2; IC5A.
COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN; PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;
                                                                                                                                                                                                                                                                                                                                                                                                                                             INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF POLYMORPH-NUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Y -> L (IN DEFECTIVE VARIANT C5D). MISSING (IN DEFECTIVE VARIANT C5D)
                                                                               MUS MUSCULUS (MOUSE).
EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
EUTHERIA; RODENTIA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               COMPLEMENT C5 ALPHA CHAIN.
                         01-NOV-1990 (REL. 16, LAST SEQUENCE UPDATE)
01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       COMPLEMENT C5.
COMPLEMENT C5 BETA CHAIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          C5A ANAPYIATOXIN.
C5B (ALPHA').
ANAPHYIATOXIN-LIKE.
BY SIMILARITY.
BY SIMILARITY.
BY SIMILARITY.
                                                                                                                                                 MEDIJNE; 90153853.
WETSEL R.A., FLEISCHER D.T., HAVILAND D.L.; J. BIOL. CHEM. 265:2435-2440(1990).
[2]
SEQUENCE OF 41-1680 FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          POTENTIAL. POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    POTENTIAL
               CREATED)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               EMBL; M35525; G309124; -. EMBL; M35526; G309123; -.
                                                                                                                                                                                                                                                                                                                     COMPLEX IS ASSEMBLED.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         18
674
674
678
678
1680
755
736
736
737
736
737
737
736
735
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1119
1633
216
1680
              90
                                                                                                                            [1]
SEQUENCE FROM N.A.
                                                                                                                                                                                                                      87185363
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            679
679
702
702
703
              01-JAN-1988
                                                                                                                                                                                                                                                                                                                                                                                          CHAIN)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DISULF ID
CARBOHYD
CARBOHYD
                                                                                                                                                                                                                         MEDLINE;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  CHAIN
PROPEP
CHAIN
PEPTIDE
CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CARBOHYD
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CARBOHYD
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SIGNAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SIGNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   /ARIANT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                VARIANT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DOMAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                      ++
                                                                                                                                                                                                                                                                                                                                   -
```

င္တ

염

음

9 9 9 9

ò

g

요

Š

용

8

US-08-487-283A-2.rsp

r-,

| SQ SEQUENCE | 1680 AA; 188877 MM; AA17044B CRC32; | à | 837 B | EQIQLIKG |
|--|--|----------------|---------------------------------------|--------------------------------|
| Query Match | 81.3%; Score 9789; DB 2; Length 1680; | ନ୍ଧ | 901 t | tftllph |
| Best Local Similarity Matches 1307; Conse | 77.8%; rvative 2 | ð | 897 T | TFTVLPL |
|)b 1 mglwgi 2y 1 MGLLGI | mglwgilcllifldktwggeqtyvisapkilrvgssenvviqvhgyteafdatlslksyp 60 | ob Qy | 961 k 957 K | kefpyri KEFPYRI |
|)b 61 dkkvtf | dkvtfssgyvnlspenkfgnaalltlgpngvpreespvshvylevvskhfskskkipit 120 | QQ | 1021 p | pvfyvfh |
| y 61 DKKFSY | SSGHVHLSSENKFQNSALLTIQFKQLPGGQNPVSYVTEVVSKHFSKSKRMPIT 120 | ď | 1017 P | PVFYVFH |
| b 121 ynngil | ynngilfihtdkpvytpdqsvkirvyslgddlkpakretvltfidpegsevdiveendyt 180 | qq | 1081 w | wltafal |
| y 121 YDNGFL | 121 YDNGFLEIHTDKPVYTPDQSVKVRVYSLADDLKPAKRETVLTFIDPEGSEVDAVEEIDHI 180 | Qy | 1077 WLTAFAL | LTAFAL |
| b 181 giisfp | giisfpdfkipsnpkygvwtikanykkdftttgtayfeikeyvlprfsvsielertfigy 240 | qq | 1141 a | agektly. |
| y 181 GIISEP | GIISFPDFKIPSNPRYGMMTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFIGY 240 | δ | 1137 AI | ARENSLY |
| 241 | knfknfeitvkaryfynkvypdaevyaffglredikdeekqmmhkatqaaklyddyaqis 300 | ය ස | 1201 101 | lgdrthpi |
| ; | | 3 | | 2014 |
|)b 301 fdseta y 301 FDSETA | fdsetavkelsynsledlnnkylyiavtvtessggfseeaeipgvkyvlspytlnlvatp 360 | 9 6 | 1261 au : 1257 T | asikikdi : : TSINIKD |
| 361 1fvkpg | Ivkpgipfsikaqvkdsleqavgypvtlmaqtvdvnqetsdletkrsithdtdgvavf 420 | qq | 1321 y | ykhegdfi |
| y 361 LFLKPG | PSKSVTRVI | δ | 1317 YI | - 5 |
| b 421 vlnlps: | vlnlpsnvtvlkfeirtddpelpeenqaskeyeavaysslsqsyiyiawtenykpmlvge 480 ::: : : | qu | 1381 fy | fylkidto |
| y 421 VINLPS | REG | ٥y | 1377 FYLKIDT | rikidt(|
| b 481 ylnimvi | ylnimvtpkspyidkithynylilskgkivgygtreklfsstyqninipvtqnmvpsarl 540 | qq | 1440 ee | eedlral |
| y 481 HLNIIV | 481 HLAIIVTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPSSRL 540 | δ | 1436 EI | EEDLKAL |
| b 541 lvyyiv | lvyyivtgeqtaelvadavwinieekcgnglqvhlspdeyvyspgqtvsldmvteadswv 600 | qq | 1500 yeyhrpdl | eyhrpdl |
| ly 541 LVYYIV | LVYYIVTGEQTAELVSDSVWIMIEEKCGNQLQVHLSPDADAYSPGQTVSLNMATGMDSWV 600 | δ | 1496 YI | YEYHRPDI |
| b 601 alsavd | alsavdravykvggnakramgrvfgaldeksdlgcgaggghdnadvfhlagltfltnana 660 : :::: | qq | 1560 et | etayaykı |
| y 601 ALAAVD | SAVYGVQRGAKKPLERVFQF1,-EKSDLGCGAGGCIANANVFHLAGLTFLTNANA 659 | δ | 1556 E | EIAYAYK |
| b 661 ddshyrd | ddshyrddsckeilrskrnlhllrgkieegaakykhsvpkkccydgarvnfyetceerva 720 | ନ୍ଧ | 1619 q | qylimgke |
| iy 660 ddsqeni | DDSQENDEPCKEILRPRRTLQK-KIEEIAAKYKHSVVKKCCYDGACVNNDETCEQRAA 716 | δ | 1616 0 | QYLIMGKE |
| b 721 rvtigp | rvtigplcirafnecctiankirkesphkpvqlgrihiktllpvmkadirsyfpeswlwe 780 | 쇰 | 1679 c | 1679 |
| ly 717 RISLGPI | RISLGPRCIKAFTECCVVASQLRANISHKDMQLGRLHMKTLLPVSKPEIRSYFPESWLWE 776 | Qy | 1676 C | c 1676 |
| b 781 ihrvpk: : : !y 777 VHLVPRI | <pre>ihrvpkrkqlqvtlpdalttweiggigiadngicvadtlkakvfkevflemnippsvvrg 840 : : : </pre> | RESULT ID C | ILT 3 CO3_NAJNA | NA |
|)b 841 eqiqlkg | eqiqlkgtvynymtsgtkfcvkmsavegictsgssaaslhtsrpsrcvfqriegssshlv 900 | AC DT | Q01833; 01-JUL-1993 01-JUL-1993 | -1993 -1993 |
| | | | | |

1376 1080 1076 1140 1136 1200 1196 1260 1256 1380 1439 1435 1499 1559 1495 1555 1618 1615 926 960 GTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKSSKCVRQKVEGSSSHLV 896 yltafsvigirkavdicptmkihtaldkadsfllentlpskstftlaivayals prfrlivsalrkeafvkgdppjyrywrdtlkrpdssvpssgtagmvettayall tqdieasshfr-lsdsgfkrijacasykpskeestsgsshavmdislptgigan cvritsateenvfvkytatllvtyktgea-adensevtfikkmsctnanlvkgk Ė 26, CREATED)
26, LAST SEQUENCE UPDATE) 1651 PRT; STANDARD; (REL.

9

Jan 28 12:73

O

SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG RESIDUES, FORMING TWO CHAINS, BETA 6 ALPHA, LINKED BY A DISULFIDE BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN, RELEASING C3A ANAPHYLATOXIN 6 GENERATING C3B (BETA CHAIN + ALPHA' FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C3, C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS. AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES. PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; REPTILIA; FRITZINGER D.C., CONNELLY M., PETRELLA E.C., BREDEHORST R., C3A ANAPHYIATOXIN.
C3B (ALPHA' CHAIN).
CLEAVAGE (BY C3 CONVERTASE).
ANAPHYLATOXIN-LIKE. COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA BASOPHILIC LEUKOCYTES. SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. INTERCHAIN (BY SIMILARITY) C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN) SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN. 2958575F CRC32; 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) COMPLEMENT C3. SIMILARITY. SIMILARITY. SIMILARITY ALPHA CHAIN BETA CHAIN. PROSITE; PS00477; ALPHA 2 MACROGLOBULIN. ₩. BY IMMUNOL. 149:3554-3562(1992) INFLAMMATORY RESPONSE; SIGNAL. 184926 L02365; G213373; -. NAJA NAJA (INDIAN COBRA) LEPIDOSAURIA; SERPENTES 1651 655 1651 738 1651 739 718 1446 1499 1578 1649 1002 1477 PIR; A46513; A46513. 1506 157 1525 164 999 100 1651 AA; HSSP; P01032; 1C5A. SEQUENCE FROM N.A. MEDLINE; 93056528 863 1346 23 23 661 661 739 738 683 684 1494 697 FISSUE=LIVER C.W. COMPLEMENT SITE DOMAIN DISULFID DISULFID DISULFID DISULFID THIOLEST DISULFID DISULFID DISULFID DISULFID DISULFID DISULFID DISULFID DISULFID CHAIN CHAIN PEPTIDE CHAIN SIGNAL CHAIN

Query Match 19.6%; Score 2367; DB 2; Length 1651; Best Local Similarity. 30.2%; Pred. No. 0.00c+00; Matches 518; Conservative 416; Mismatches 655; Indels 127; Gaps 97;

4 malylvaalligfpg-s-shgalytlitpavlrtdteegilveahgdstpksldifvhdf

셤

60 PDKKESYSSGHVHLS-SENKFQNSAILTIQPKQLPGCQNPVSYYYLEVVSKHFSKSKRMP 118 121 lsyqsgfvfiqtdkgiytpgspvryrvfsvdhnmhrmdktvivefqtpegivvs-skpvn 179 237 292 62 prkqktlfqsrvdmnqagsmfvtpti-kvpakelnkdskqnqyvvvkvtgpqvalekvvl 120 293 eatlkrdt-lrs-rfqdlnqlvghtlyvsvtvitesgsdmvvteqggihivtspyqiyft 350 397 417 398 aklilntplnigslpitvrtnhgdlprergaiksmtatayqtqggsenylhvaitsteik 457 517 536 PSSRLLVYYIVTGEQTAELVSDSVWLNIEEKCGNQLQVH-LSPDADAY-SPGQTVSLNWA 593 634 694 qry-r-qqfpikalssravpfvivpleqglhdveviasvrgelasdgvrkklkvvpeger 927 TKSSKCVRQKVEGSSSHLVTFTVLPLEIGLHNINFSLETR--FGKEILVKTLRVVPEGVK 935 518 psfrfvayyqv-gnn--eivadsvwvdvkdtcmgtlvvkgassrddriqkpgaamkikle 574 LINANADDSQENDEPCKE-ILRPRRT---LQKKIEEIAAKYKHSVVKKCCYDGACVNNDE 709 710 -TCEQRAARISLGPRCIKAFTECCVVASQLRA-NISHKDMQLGRLHMKTLLPVSKPEI-R 766 sdfpeswlwlteeltgepnnggissktvpfylrdsittwellavglsptkgicvaepyei 814 767 SYFPESWLMEVH-LV--PR-R----KQLQFALPDSLTTWEIQGIGISNT-GICVADTVKA 817 815 tvmkdffidlrlpysvvknegveirailynyadedi-y-vr---veliynpafcsasteg 869 877 928 knivtiieldpsvkgv-ggtqeltviankld-dkvpdtevetrisvlgdpvaqiiensid 985 695 ytcekrakyıqegdackaaflecchyikgirdenqreselflarsdfedelfgddniisr 754 psgsir-p-ynlpelvsfgtwkavakyehspeesytayfdvreyvlpsfevrlqpsdkfl -yidgnknfhvsitarylygkkv-egvafvvfgvk--i-ddakksipdsltripiidgdg 351 ktpkyfkpgmpyeltvyvtnp-d---gs-paa-hvp-v-vs-ea--ih-segtt-lsdgt pgdnlpvnfnvrgnanslnqikyftylilnkgkifkvgrqprrdgqnlvtmnlhitpdli gdpgarvglvavdkavyvlndkykisqakiwdtieksdfgctagsgqnnlgvfedaglal TOMDSWVALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGLNNANVFHLAGLTF ttstnlntkgrsaakcpgpanrrrrssvllldskaskaaqfqdqglrkccedgmhenpmg KVFKDVFLEMNIPYSVVRGEQIQLKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQG :: -: = = = = : : : : : ::: ==::: .. = = = 180 458 654 755 818 238 575 870 878 원 പ്പ õ 셤 g õ a 음 셤 셤 õ ð ð ò ð ò ð ð ð g ð 셤 ð 셤 ð В ò Р ð g ð 9

음

ð

용

8

Jan 28 12:23

NETTESHEIM D.G., EDALJI R.P., MOLLISON K.W., GREER J., ZUIDERWEG E.R.; PROC. NATL. ACAD. SCI. U.S.A. 85:5036-5040(1988). MATANABE Y., MATSUI N., YAN K., NISHIMUKAI H., TOKUNAGA K.,
JUJI T., KOBAYASHI N., KOHSAKA T.;
MOL. IMMUNOL. 30:62-6211993).
-!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL
REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.
AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE BOTTO M., YONG FONG K., SO A.K., KOCH C., WALPORT M.J.; J. EXP. MED. 172:1011-1017(1990). SINGER L., WHITEHEAD W.T., AKAMA H., KATZ Y., FISHELSON MEDLINE; 82174534. THOMAS M.L., JANATOVA J., GRAY W.R., TACK B.F.; PROC. NATL. ACAD. SCI. U.S.A. 79:1054-1058(1982) PROC. NATL. ACAD. SCI. U.S.A. 82:708-712(1985) Р. Л. LACHMANN P.J.; LAMBRIS J.D.; LACHMANN ISAAC L., ISENMAN D.E.; J. BIOL. CHEM. 267:10062-10069(1992). ERRATUM (RETRACTION OF ABOVE ARTICLE) MUTAGENESIS OF THIOESTER BOND REGION BIOL. CHEM. 269:28494-28499 (1994) J. BIOL. CHEM. 250:8293-8301(1975) SEQUENCE OF 1409-1563.
MEDLINE, 88154452.
DAOUDAKI M.E., BECHERER J.D., L
J. IMMUNOL. 140:1577-1580(1988) POZNANSKY M.C., CLISSOLD P.M., J. IMMUNOL. 143:1254-1258(1989) POZNANSKY M.C., CLISSOLD P.M., J. IMMUNOL. 143:3860-3862(1989) DOLMER K., SOTTRUP-JENSEN L.; FEBS LETT. 315:85-90(1993). VARIANTS GLY-102 AND PRO-314. BRUIJN M.H.L., FEY G.H.; STRUCTURE BY NMR OF C3A SEQUENCE OF 988-1036, SEQUENCE OF 672-748. MEDLINE; 88276894. VARIANT C3F/S. MEDLINE; 89309808. MEDLINE; 76069169. MEDLINE; 85140166. MEDLINE; 93106233. SEQUENCE FROM N.A MEDLINE; 92250565 MEDLINE; 91011240. MEDLINE; 95050640. MEDLINE; 90063087 VARIANT GLN-1320 DISULFIDE BONDS VARIANT ASN-549. WETSEL R.A.; HUGLI T.E.; 6 qqpdqvfkenapvihgemlggtkgaep-easl--tafivtallesrsvckeqinildssi 1158 1318 tvsas-gdgkatmtiltvynaqlredanvcnkfhldvsvenvelnlkgakggkaalrlki 1376 986 gsklnhliitpsgcgeqnmitmtpsviaty-yldatgqwenlgvdrrte-a-ikqimt-g 1041 1042 yaqqmvykkadhsyaaftnrassswltayvvkvlamasnmvkdisheiicggvkwlilnr 1101 1053 MISIMSYRNADYSYSVAWKGGSASTWLTAFALRVLGQVNKYV-EQNQNSICNSLIALVENY 1111 1159 nkatdylikkyekiqrpyttaltayalaaadrindd-r--v--lm-aa-stgrn---r-w 1207 -e--ey-narthn----iegtsyallallkmkkfaevgpvvrwlidqkyyggtygqtq 1257 1258 atvmvfqalaeyeiqmpthqdlnldisiklperevperysindrnavqartvetklnedf 1317 1492 kichgnicrcaeetcsllngq-k-kidlg1rigkacagnvdyvyktk1lrieekdgndiy 1549 1432 vviyldkvshsedeclhfkihkhfevgfiqpgsvkvysyynldeqctkfyhpdketglln 1491 1518 KVCEGAACKCVEADCGOMQEELDLTISAETRKOTACKPEIAYAYKVSITSITVENVFVKY 1577 1550 fmdvlevikggtdrnagakarqyvsgrkcqealnlkldndyliwglssdlwpmk-d-dis 1607 :: :: ||| :|| :|: |
936 RESYSCYTLDP—RGIYGTISRRKEFPYRIPLDLVPKTEIKRILSVKGLLVGEILSAVLS 993 EUKARYOTA; METAZOA; CHORDATA; VENTEBRATA; TETRAPODA; MAMMALIA; EUTHERIA; PRIMATES. [1] 21-JUL-1986 (REL. 01, CREATED)
21-JUL-1986 (REL. 01, LAST SEQUENCE UPDATE)
01-OCT-1996 (REL. 34, LAST ANNOTATION UPDATE)
COMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN). 1608 ylit-kn-twierwpnedecqd-eef-qnlcddfag 1639 1635 YIYPLDSLTWIEYWPRDTTCSSCQAFLANL-DEFAE 1669 PRT; 1663 AA STANDARD; HOMO SAPIENS (HUMAN)

1208

g

ð

유

ð

\$

요

ð g ð 연 \$ ብ à g ð

1102

ð

ð

g

õ

_ =

LT 4 CO3_HUMAN

g

ð

RESULT
1D CO
AC PO
DT 21
DT 21
DT 01
OE CO
GN C3
OS HG
OC EU
RN [1]

| ٠ | | | | | | |
|----|-----|----|----|---|----|---|
| | ю | × | х | × | | |
| ۰ | 8 | 88 | ٠. | ۰ | × | ٠ |
| х | × | × | ۰ | х | ۰ | |
| ۰ | 30 | 84 | × | ۰ | м | |
| ۰ | × | • | × | ۰ | × | • |
| к | - | 80 | ۰ | × | ۰ | |
| ۰ | × | × | ۰ | х | ۰ | |
| ۰ | | ٠ | | ۰ | | |
| ۰ | : 6 | | н | ۰ | × | |
| к | ٠ş | 7 | v | ж | ۰ | 1 |
| s | | ч | ٠. | × | | |
| ۰ | : • | ٠ | H: | ۰ | × | |
| ۰ | | 8 | ١. | 0 | × | |
| 4 | | | ŀ | и | 0 | |
| o | ٠. | 8 | · | × | | |
| | 20 | ٠. | × | ٥ | | |
| | | × | ю | 0 | 2 | ۰ |
| • | c | 1 | ø | я | ٠, | |
| м | • | | ς. | ю | ۰ | |
| 1 | м | | м | | × | |
| ٩. | 20 | × | м | ۰ | | ۰ |
| Ł | ж | | ۲. | ۰ | ٠. | • |
| В | и | ч | Ŀ | × | ۰ | |
| т | ٠ | | ŀ | | ٠, | |
| 7 | 2 | 8 | × | ٠ | × | • |
| 2 | 0 | ٠, | О | × | | ۰ |
| ٠ | ٠ | × | ø | х | × | |
| 1 | | | | | ٠. | |
| 1 | н | S | ۲ | 0 | × | ۰ |
| 2 | - | z | ø | х | | ۰ |
| k | v | | ŗ, | × | ۰ | ١ |
| ۸, | ٠. | ı. | ۰ | | | ٠ |
| 3 | 3 | 2 | × | 0 | | ۰ |
| ı | × | o | r | ٠ | | ۰ |
| В | - | 2 | | | ٠. | • |
| к | | 0 | Ç٠ | х | | |
| | | | × | ۰ | | • |
| ٠ | ж | | х | ۰ | × | • |
| 7 | | | r. | • | ٠. | ۰ |
| ٠ | ٠ | × | ۰ | × | ۰ | • |
| | ð. | м | r: | o | | |
| 4 | 4 | • | ١. | ۰ | × | • |
| × | | | ۰ | × | ٠. | ۰ |
| х | × | × | ۰ | × | ю | |
| 8 | - | 88 | х | × | | |
| ۰ | × | • | ж | • | × | • |
| ۰ | | | | | | |
| | 20 | | | | × | |
| × | | 8 | ٥ | × | × | |
| Š | | × | Ì | ì | Š | |
| į | | | | ۱ | | |
| ۱ | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |

RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN, RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA' SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES CHAIN).

C3B IS RAPIDLY SPLIT IN TWO POSITIONS BY FACTOR I AND A COFACTOR TO FORM IC3B (INACTIVATED C3B) AND C3F WHICH IS RELEASED.

-!- IC3B IS THE SLOWING CLEANED (POSSIBLY BY FACTOR I) TO FORM C3C AND C3DG. OTHER PROTEASES PRODUCE OTHER FRAGMENTS SUCH AS C3D OR C3G. -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLÓBULIN. -!- ENCTION: DERIVED FROM PROTEOLYTIC DECRADATION OF COMPLEMENT C3, C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND

POLYMORPHISM: THERE ARE TWO ALLELES: C3S (C3 SLOW), THE MOST COMMON ALLELE IN ALL RACES AND C3F (C3 FAST), RELATIVELY FREQUENT IN CAUCASOIDS, LESS COMMON IN BLACK AMERICAN, EXTREMELY RARE IN BASOPHILIC LEUKOCYTES

DISEASE: C3 DEFICIENCY CAUSES A SUSCEPTIBILITY TO PYOGENIC ORIENTALS.

-!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN

INFECTION.

EMBL; K02765; G179665; -.
PIR; A01257; C3HU.
PIR; A27603; A27603.
HSSP; P01032; 1C5A.
SWISS-2DPAGE; P01024; HUMAN.

MIM; 120700,

INFLAMMATORY RESPONSE; GLYCOPROTEIN; SIGNAL; POLYMORPHISM; COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.

DISEASE MUTATION. SIGNAL CHAIN

C3A ANAPHYLATOXIN. C3C FRAGMENT. COMPLEMENT C3. ALPHA CHAIN. BETA CHAIN. CHAIN PEPTIDE CHAIN PEPTIDE CHAIN

C3G FRACMENT.
C3D FRACMENT.
C3F FRACMENT.
CLEAVAGE (BY C3 CONVERTASE).
CLEAVAGE (BY FACTOR I).
CLEAVAGE (BY FACTOR I).
CLEAVAGE (BY FACTOR I). 1002 1304 748 954 954 1303 1320 693 PEPTIDE PEPTIDE PEPTIDE PEPTIDE

ANAPHYLATOXIN-LIKE. PROPERDIN-BINDING INTERCHAIN DISULFID SITE SITE SITE SITE DOMAIN DOMAIN

559 627 693 694 DISULFID DISULFID DISULFID

DISULFID

CARBOHYD

ß

US-08-487-283A-2.rsp Jan 28 12 23

7

| | POTENTIAL. | | R -> G (IN ALLELE C3F). | L -> P. | D -> N (IN C3 DEFICIENCY; IMPAIRS | SECRETION). | R -> Q (IN C3 DEFICIENCY; ALLOTYPE | C3'F02'; MAY INHIBIT IC3B SYNTHESIS). | D -> N (IN REF. 2). | $E \rightarrow Q$ (IN REF. 2). | H -> S (IN REF. 4). | 3 MW; 840B4548 CRC32; | |
|----------|------------|----------|-------------------------|---------|-----------------------------------|-------------|------------------------------------|---------------------------------------|---------------------|--------------------------------|---------------------|-----------------------|--|
| 939 | 1617 | 1013 | 102 | 314 | 549 | | 1320 | | 681 | 700 | 1026 1026 | AA; 18716 | |
| 939 | 1617 | 1010 | 102 | 314 | 549 | | 1320 | | 681 | 700 | 1026 | 1663 | |
| CARBOHYD | CARBOHYD | THIOLEST | VARIANT | VARIANT | VARIANT | | VARIANT | | CONFLICT | CONFLICT | CONFLICT | SEQUENCE | |
| E | F | 占 | F | F | F | F | ij | F | FŢ | F | FI | ŏ | |

458; Mismatches 660; Indels 115; Gaps 91; Score 2361; DB 2; Length 1663; Pred. No. 0.00e+00; Best Local Similarity 28.4%; 489; Conservative Query Match Matches

64 8 sll-ll-llthlplalg-spmysiitpnilrleseetmvleahdaggdvpvtvtvhdfpg ð 65 kklvlssektvltpatnhmgnvtftipanrefksekgrnkfvtvgatfgtqvve-kvvlv 123

셤

В

Š

ð

62 KKFSYSSGHVHLSSENKFQNSAILTIQP-KQLPGGQNPVSYVYLEVV-SKHFSKSKRMPI 119

124 slqsgylfiqtdktjytpgstvlyriftvnhkllpvgrtvmvnienpegipvkqdslssq 183 184 nqlqvlpls-wdipelvnmgqwkirayyenspqqvfstefevkeyvlpsfeviveptekf 242 DHIGHISFPDFKIPSNPRYGMMTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEP-E-Y 235 임

178

ð

296 243 yyi-yne-kglevtitarflygkkv-egtafvifgi-qd--gegrislpeslkripiedg 염

236 NFIGYRNFRNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLING 295

ð

sgevvlsrkvlldgvqnlraedlvgkslyvsatvilhsgsdmvqaersgipivtspyqih 356 296 IAQVTFDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLM 355 297 g ð

ftktpkyfkpgmpfdlmvfvtnp-d---gs-pay-rvp-vav-q--ge-dtvqsltq-gd 404 357 셤

gvaklsinthpsqkplsitvrtkkqelseaeqatrtmqalpystvgnsnnylhlsvlrte 464 356 405 \$ g

à 465 lrpgetlnvnfllrmdraheakiryytylimnkgrllkagrqvrepgqdlvvlplsittd 524 g 8

591 525 fipsfrlvayytligasggrevvadsvwvdvkdscvgslvvksggsedrgpvpgggmtlk 셤 ð 644 585 iegdhgarvvlvavdkgvfvlnkknkltgskiwdvvekadigctpgsgkdyagvfsdagl

<u>.</u>

651 592 MATCMDSWVALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGLINNANVFHLAGL :: :: à 645 tftsssgqqtaqraelqcpqpaarrrrsvqltekrmdkvg-kypkel-rkccedgmrenp 702 ::

셤

| 9 | | | |
|----|-----|------------|-----|
| | 0 | | w |
| | | ** | |
| | 90 | 92 | ۰ |
| | | | × |
| | | • | × |
| 0 | | | • • |
| œ | 88 | ж. | ** |
| 88 | ٥. | ٠. | × |
| ж | | | |
| | | | ۰ |
| 0 | 33 | | |
| | ••• | • | |
| × | • | • | • |
| | • | 00 | |
| | | | |
| | ٠., | | |
| × | | | × |
| ж. | | | • |
| | œ | | |
| ж. | 88 | • | |
| 8 | | | |
| | | | • |
| w | 2 | ж. | • • |
| 8 | | 83 | 83 |
| 88 | • | ** | |
| | | | |
| • | | | |
| | | ٠., | ٠. |
| × | | | |
| 8 | œ | | |
| 0 | 2 | | • |
| | | ٠., | |
| × | | | |
| ۰ | | | |
| 0 | | | |
| | | | |
| | • | | • |
| | | | × |
| ۰ | | | |
| | | | |
| 8 | | | |
| | | • | |
| | | | |
| 8 | 33 | 80 | |
| 8 | | | |
| | | κ. | |
| × | | | |
| 0 | • | • | |
| • | 3.4 | Ψ. | |
| | | -4 | 88 |
| ж | | | |
| | ٠. | О. | |
| ø | | 80 | |
| × | 80 | | ** |
| ٠, | | - | |
| | œ | | |
| | | • | 99 |
| | | H. | ** |
| ж | и, | 12 | |
| Э, | ~ | ۰. | œ |
| | | | |
| 8 | | | |
| P | ٠. | 3 | × |
| Ì | Ċ | Ž | × |
| | | Ž | ä |
| | ٥ | Ž | |
| | Ċ | 2 | |
| | Ċ | 7 | |
| | Ċ | Š | |
| | Ċ | 2 | |
| | • | 2 | |
| | | | |
| | | 2 | |
| 1 | | | |
| | | 2 | |
| | | 3 | |
| | | | |
| | | | |
| | | | |
| • | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | 7 3 3 | |
| | | 7 3 3 | |
| | | | |
| | | 3.3 | |
| | | 3.3 | |
| | | 3.3 | |
| | | | |
| | | 73.3 | |
| | | 3.3 | |
| | | | |
| | | 3.3 | |
| | | 3.3 | |
| | | | |
| | | 3.3 | |
| | | 3.3 | |
| | | 3.3 | |
| | | | |
| | | | |
| | | 7.073.33 | |
| | | 7.073.33 | |
| | | 7.73 | |
| | | 7.073 | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | 7-10-00-50 | |

US-08-487-283A-2.rsp Jan 28 12 23

9

1578 KATLIDIYKTGEAVAEXDSEITFIKKVTCTNA-ELVKGRQYLIMGKEALQIKYNFSFRYI 1636 ò

1661 1620 yiigkdtwvehwpeedecqdeenqkqcqdlgaftesmvvfgc 유

| : ||:| ||: | : : : || 157 | 1676 | 1637 YPLDSLTWIEYMPRDTTCSSCOAF-L-ANLDEFAEDIFINGC 1676

ð

FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. AFTER ACTIVATION (C3B), IT CAN BIND COVALENTLY, VIA ITS REACTIVE THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES. CYCLOSTOWATES C3 APPEARS TO REPRESENT THE COMMON ANCESTOR OF MAMMALIAN C3 AND C4, SHOWING SIMILARITIES TO ALPHA CHAIN (BY SIMILARITY).
GAMMA CHAIN (BY SIMILARITY).
C3A ANAPHYLATOXIN (BY SIMILARITY).
ANAPHYLATOXIN-LIKE.
BY SIMILARITY.
BY SIMILARITY.
BY SIMILARITY.
BY SIMILARITY.
BY SIMILARITY.
SY SIMILARITY.
BY SIMILARITY. 01-FEB-1996 (REL. 33, CREATED)
01-FEB-1996 (REL. 33, LAST SEQUENCE UPDATE)
01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
COMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN) (FRAGMENT) COMPLEMENT PATHWAY; PLASMA; INFLAMMATORY RESPONSE; GLYCOPROTEIN; !- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN. !- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. COMPLEMENT C3. BETA CHAIN (BY SIMILARITY) LAMPETRA JAPONICA (JAPANESE LAMPREY). EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; PISCES; AGNATHA (CYCLOSTOMATA). PRT; 1673 AA. POTENTIAL. J. IMMUNOL. 148:3290-3295 (1992). 187767 MEDLINE; 92251197. NONAKA M., TAKAHASHI M.; STANDARD; 1673 653 1375 1375 1673 713 705 712 713 33,33 EMBL; D10087; G222888; 692 71 986 99 1673 AA; [1] SEQUENCE FROM N.A. BOTH PROTEINS. 1 <1 14 14 657 657 657 657 678 678 TISSUE=LIVER LT 5 CO3_LAMJA DISULFID DISULFID THIOLEST SEQUENCE CHAIN CHAIN PEPTIDE DOMAIN DISULFID Query Match NON TER SIGNAL SIGNAL. CHAIN

Gaps 105; 458; Mismatches 636; Indels 130; Score 2309; DB 2; Length 1673; Pred. No. 0.00e+00; 19.2%; 29.5%; 512; Conservative Local Similarity Best Loc Matches

59 latqtitltqldpaiatlklgfdienpdktnsastkhhvrlv-akvesksfnkeitahal 117 65 1 vl-llmsvqtsvtqdpmvllsvpsviligsdvnvlvdhaastedvsvvvraeefltkk-q 58 6 ILCFLIFLGKTWGQEQTYVISAPKIFRVGASENIVIQVYGYTEAFDATISIKSYPDKKFS 유 g ð

66 YSSGHVHLSS-ENKFQNSAI-LTIQ-P-KQLPGG-QNPVSYVYLEVVSKHFSKS-K-RMP 118 g õ

119 8

용 ð 셤 Š g Š g Š a ð g õ 유 Š 요 Š g Š පු

| 178 ktikatdegivdgtsftipaiskhgtwkifarmsgapninssaefdvreyilptfevkin 237 - :- :-:: : 175 EEI-D-HIGIISFPDFKIPSNPRYGMYTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIE 232 | Db Qy | 1092 gs: |
|--|---------------------------------------|--|
| 238 pkgrvfhind-eefvvditanyfnqelvsgta-yvryfl-en-gdvpk-lvdsssttl 290 :::: : :: :: 233 PEYNFIGYKNFKNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTM1, 292 | DP Qy | 1150 fl. |
| 291 vageglsilkkekllk-l-fpnakdllafsltikttvlssgaaeteeaelvgikivesry 348 : : : : : : : : | d ç | 1206 e- : 1233 QHI |
| 349 qitatktsryfkpelpyfiqvevrnadgspskevdvvak-vqvgsatinpqk-m-r 401 : :: : : : : : | do yo | 1263 ald |
| 402 tdsngltsftvt-ppnvngltvtvrtderh-psneggelvytaqkyas-a-symhid-v- 455 : :: : : : :: : : : : | a & | 1321 kau : 1349 ST |
| 456 t-rimrlgetlnvfltakttqlnavthftymvltrgvivktnrktkesgggpsnvripit 514 ::: ::: : : ::: : :: : ::: ::: : 472 NHKALLVGEHLMIVTPKSPYIDKITHYNYLLLSKGKIIHFGTREKFSDASYQSINIPVT 531 | da vo | 1380 dio |
| <pre>515 pdmaprfrflayyilpggeivadsvtvevtelcksqvslslkgrp-tlepkamltld 570 : : : : : : ::: : : 532 QNWVPSSRLLVYYIVTGEQTAELVSDSVWINIEEKCGNQLQVHLSPDADAYSPGQTVSLN 591</pre> | g & | 1440 yiu : 1449 LE |
| 571 ligepdarvgllavdqavyavnrkhrltqdrvwkametfdtgctaeggagrpgvfsdagl 630 : :: : : : : | g & | 1500 yn. : 1509 YS |
| 631 alitskglnttdrseigcpkvpsrkprqlsmlqirreaekytqef-rkccvdglkmsptg 689 ::: : : :: :: : : : 652 TFLTNANADDSQENDEPCKEIL-RPRRTLQK-KIEEJAAKYKHSVVKKCCYDGACVN-ND 708 | a v | 1559 dni 1567 SE |
| 690 ggceerlkrut-gpkecvdaflgcckkaeeyrkseslgaktulrindfmeldimnedevn 748 : : :: : : | අ ර | 1618 kg- : 1623 EAI |
| 749 mmayfpqswqwnkyknsckygrhpqirlqlpdtittwnmqavsisktrgvcladplllvs 808 :: : : :: :: :: :: : | RES ID AC DT | RESULT 6 ID CO3 MOUS AC PO1027; DT 21-JUL-1 |
| | E E E | 01-FEB-1 01-FEB-1 COMPLEME C3. |
| SSKCVRQKVEGSSSHLVTFTVLPLEIGLANINF-S-LETW-FGKEILVKTLRVVPEGVKR | 00 00 00 | MUS MUSC EUKARYOT EUTHERIT |
| 920 di-r-s-esrsvh-veeretffikneispdvvpnsdvltfisvkgdelaetmvncldak 974 : ::::: : ::: : - 937 ESYSCYTLDPRGIYGTISRRKEF-PYRIPLDLVPKTEIKRILSVKGLLVGEILSAVLSQE 995 | R R 8 | (1) SEQUENCE MEDLINE; |
| 975 sisnliqiptgqqqqmmikmapttltliyldsvqeweki-qlhrreeaiqflkqqysr 1031 | 2 2 2 2 2 | DOMDEY B PHILOS. [2] |
| 1032 elsyrkadhsyaafikrpsstwltafvvkvyslakrviivdnqelcgpvewiiknrqmsd 1091 | # # # # # # # # # # # # # # # # # # # | MEDLINE; WETSEL F J. BIOL. |

유

Š

9

ð

셤

δ

g

Š

g

õ

원 장

õ

1379 laskvsdlkrmytiaitryalalqd---peseaahsswkklenrttfeskghrywkae- 1205 1262 isnyeitdsvliiywdkvpstedycfafkikqmlrsdmiqpvtasvydyyspadkctrl 1499 lgamagykmdssskel-idvqleit-spknnfekkmkiteetrfvgephkippggniti 1320 IECLTEYSL-LVKQLRLSMDIDVSYKHKGALHN-YKMTDKN-FLGRPVEVLLNDDLIV 1348 :::|:| |: :|: : :||:: | |:| ::: : | |:|| | || :|| ETDYQIKDCHVILQLASIPSSDFLCVRERIFELFEVGFLSPATFTVYEYHRPDKQCTMF 1508 nrtevgsfvyytvniqtviksgqdqaiqpkatrl-fivtrscdgrlgmetprqyllmgr 1617 ALQIKYNFSFRYIYPLDSLTWIEYWPRDTTCS-SCQAFLANLDEFAEDIFLNGC 1676 -t-shvlrmsais-veatayglltylrkkdyesareivdwltegrnygggfqstqdtil as-grgtftlsimsvfnkvapsskscstfdlkvtmteaddgespqgrlgwfdgkrrrrr g-etk-drndrfgyvldasswvegwpvdekcngpnvqtfcaikreyefsmqiqgc 1671

EBULT 6

CO3 MOUSE STANDARD; PRT; 1663 AA.

C CO3 MOUSE STANDARD; PRT; 1663 AA.

T 21-JUL-1986 (REL. 01, CREATED)

T 01-FEB-1996 (REL. 33, IAST SEQUENCE UPDATE)

T 01-FEB-1996 (REL. 33, IAST SEQUENCE UPDATE)

E COMPLEMENT G3 PRECURSOR (HSE-MSF) (CONTAINS: C3A ANAPHYLATOXIN).

C 3.

MUS MUSCULUS (MOUSE).

C EUTHERIA; RODENTIA.

C EUTHERIA; RODENTIA.

I 11

P SEQUENCE FROM N.A.

A MEDLINE; 85038864.

A MEDLINE; 85038864.

A DOMDEY H.;

D PHILOS. TRANS. R. SOC. LOND., B, BIOL. SCI. 306:333-344(1984).

I 21

C MEDLINE; 85054819.

A METSEL R.A., LUNDWALL A., DAVIDSON F., GIBSON T., TACK B.F., FEY G.H.;

J. BIOL. CHEM. 259:13857-13862(1984).

| 13 SEQUENCE OF 671-748 FROM N.A. DOMENT H., WIEBBURS K., KRZAMIER H., WILLER V., ODINK K., FEY C.H.; PROC. MAIL. ACAD. SCI. U.S.A. 79:1619-7623(1992). SEQUENCE OF 671-748 FROM N.A. PROC. MAIL. ACAD. SCI. U.S.A. 79:1619-7623(1992). SEQUENCE OF 1-34 FROM N.A. PROC. MAIL. ACAD. SCI. 47:300-312(1993). SEQUENCE OF 1-34 FROM N.A. SEQUENCE OF 1-34 FROM N.A. SEQUENCE OF 2-44 AND 149-760. WEDLINE, 9131334. AND SEQUENCE OF 1-34 FROM N.A. SEQUENCE OF 2-44 AND 149-760. AND SEQUENCE OF 2-44 AND 149-760. AND SEQUENCE OF 2-44 AND 149-760. AND SEGUENCE OF 2-44 AND 149-760. AND S | .A. | ZMAIER M., MULLER V., ODINK K., FEY G.H.; .A. 79:7619-7623(1982). | .e. | DEV H. 3 | 07-312 (1983) . | | | GGELMANN H., FEY G.; .A. 79:7077-7081(1982). | 760. | | G., MIKI K., NICOLSON G.L.; | 593). ENTRAL ROLE IN THE ACTIVATION OF THE | PROCESSING BY C3 CONVERTASE IS THE CENTRAL | ICAL AND ALTERNATIVE COMPLEMENT PATHWAYS. AN BIND COVALENTLY. VIA ITS REACTIVE | RFACE CARBOHYDRATES OR IMMUNE AGGREGATES. | IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG CHAINS, BETA 6 ALPHA, LINKED BY A DISULFIDE | TIVATES C3 BY CLEAVING THE ALPHA CHAIN, | TOXIN & GENERATING C3B (BETA CHAIN + ALPHA' | C3B IS RAPIDLY SPLIT IN TWO POSITIONS BY FACTOR I AND A COFACTOR | TO FORM ICSB (INACTIVATED C38) AND C3F WHICH IS RELEASED. | AVED (POSSIBLI BI FACTOR 1) TO FORM CSC AND PRODUCE OTHER FRAGNENTS SUCH AS C3D OR C3G. | AND ALPHA-2-MACROGLOBULIN. | PROTECLYTIC DEGRADATION OF COMPLEMENT C3, MEDIATOR OF LOCAL INFLAMMATORY DROCFS IT | N OF SMOOTH MUSCLE, INCREASES VASCULAR | S HISTAMINE RELEASE FROM MAST CELLS AND | NE ANAPHYLATOXIN-LIKE DOMAIN. | | | | MACROGLOBULIN. | MENT ALTERNATE PATHWAY; PLASMA; | COPROTEIN; SIGNAL. | COMPLEMENT C3. | BETA CHAIN. | C3A ANAPHYLATOXIN. | C3B (ALPHA' CHAIN). | C3G FRAGMENT. | C3G FRAGMENT. | C3D FRAGMENT. | | CLEAVAGE (BY FACTOR I). |
|--|--|---|--------------------------------------|--------------------|--------------------------------------|-----------------------------------|--------------------|--|---------------------------------------|--------------------|---|---|--|---|---|---|---|---|--|---|--|--------------------------------------|--|--|---|-------------------------------|------------------------|----------------------|--------------------|---|---|--|----------------|-------------|--------------------|---------------------|---------------|---------------|---------------|---------|-------------------------|
| | (3) SEQUENCE OF 671-748 FROM N.A. MEDLINE: 83117730. | DOMDEY H., WIEBAUER K., KAZMAIER M. PROC. NATL. ACAD. SCI. U.S.A. 79:76 | [4] SEQUENCE OF 671-748 FROM N.A. | MEDLINE; 84201365. | ANN. N.Y. ACAD. SCI. 421:307-312(19) | (5) SEQUENCE OF 1-34 FROM N.A. | 4EDLINE; 83117622. | WIEBAUER K., DOMDEY H., DIGGELMANN H., FEY G.; PROC. NATL. ACAD. SCI. U.S.A. 79:7077-7081(1982) | [6] SEQUENCE OF 25-41 AND 749-760. | AEDLINE; 93373334. | HAMADA JI., CAVANAUGH P.G., MIKI PANGED DES 53.4439.443341993 | .ANCER RES. 33:4418-4423(1993). -!- FUNCTION: C3 PLAYS A CENTRAL ROI | COMPLEMENT SYSTEM, ITS PROCESSIN | REACTION IN BOTH CLASSICAL AND A AFTER ACTIVATION C3B CAN BIND CO | THIOLESTER, TO CELL SURFACE CARI | -!- SUBUNIT: C3 PRECURSOR IS FIRST I RESIDUES, FORMING TWO CHAINS, BI | BOND. C3 CONVERTASE ACTIVATES C | RELEASING C3A ANAPHYLATOXIN & GI | | | | -!- SIMILARITY: TO C4, C5 AND ALPHA- | | INDUCES THE CONTRACTION OF SMOOT | PERMEABILITY AND CAUSES HISTAMIN | | EMBL; K02782; G309122; | MBL; M33032; E22244; | VIR; A05290; C3MS. | 133F; FULUSZ; ICJA: PROSITE: PS00477; ALPHA 2 MACROGLOBI | COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA; | INFLAMMATORI RESPONSE; GLICOPROTEIN; SIGNAL 1 | 25 1663 | 25 666 | E 671 748 | 749 1663 | 955 1303 | 955 1001 | 1002 1303 | 748 749 | 1303 1304 |

| SIMILARITY). | tch 19.1%; Score 2296; DB 2; Length 1663; al Similarity 28.2%; Pred. No. 0.00e+00; 479; Conservative 447; Mismatches 665; Indels 106; Gaps 84; ysiitpnvlrleseetivleahdaggdipvtvtvqdflkrqvltsektvltgasghlrsv 87 | 14 14 20 | VRVYSLNDDIKPAKRETVLTFID-PEGSEVDMYEEIDHIGIISFPDFKIPSNPRYGMM 199 kirafyehapkqifsaefevkeyvlpsfevrveptetfy-yiddpnglevsiiakflygk 264 : : | <pre>vgkslyvsvtvilhsgsdmveaersgipivtspyqihftktpkffkpampfdlmvfvtnp 380</pre> | asggrevv 54 : EQTA_ELV 55; gvfvlnkk 60; : ::: AVYGVQRG 61; |
|--|--|--|---|---|---|
| ANAPHILATOXIN-L INTERCHAIN (BY BY SIMILARITY. | ; Score 2296; DB 2; ; Pred. No. 0.00e+00; 447; Mismatches 665 ahdaqqdipvt.vt.qdflkrqv: ; :: : : | tvvanfgetvoekavmvsfi ::::: YLEVVSKHFSKSRRMPITYI Lietpdgipvkrdilssnn :::::::::::::::::::::::::::::::::: | <pre>rVLTFID-PEGSEVDMVEEID) sfevkeyvlpsfevrveptetfy- i </pre> | eaersgipivtspyqihftkt : ::: EALIPGIKYULSPYKLALVAT := akaltq-ddgvak := -: - QETSDLDPSKSVTRVDDGVAS tmhnsnnylhlsvsrmelkpg :: | epgqdlvvlslpitpefij :: : : : : EEDASYQSINIPVTQNMVT prdnhlapgqqttlriegr EPADAYSPGQTVSLNMATC spagykmagyfmdaglafkt |
| NALIN 693 728 SULFID 559 816 SULFID 626 661 SULFID 693 720 SULFID 693 720 SULFID 694 727 SULFID 101 1158 SULFID 1101 1158 SULFID 1389 1458 SULFID 1506 1511 SULFID 1506 1511 SULFID 1508 1458 SULFID 1508 1458 SULFID 1509 1458 SULFID 1518 1590 SULFID 1537 1661 SULFID 1637 1637 SULFID 1637 1637 | 19.1%; ocal Similarity 28.2%; s 479; Conservative g ysiltpnvlrleseetivlea | sikipaskefnsdkeghkyv | VRVYSLNDDLKPAKRETVLTFID-PEGSEVDMVEEIDHIGIISFPDFKIPSNP kirafyehapkqifsaefevkeyvlpsfevrveptetfy-yiddpnglevsiiak : | <pre>vgkslyvsvtvilhsgsdmveaersgipivtspygihftktpkffkpampfd</pre> | ryytylwmnkgkllkagrqvi :: :: THYNYLILSKGKIIHFGTREF adsvwvdvkdscigtlvvkgc : ::: SDSVWLNIEEKCGNQLQVHLS |
| DOMAIN DISULEID THOLEST EQUENCE | Query Match Best Local Matches (28 ys) | 88 83 148 | 143 206 200 265 259 | 321 319 381 379 428 | |
| | Qu Be Ma Ma Ma Ma Ma Ma Ob | d by d | cy by cy | 4 6 6 6 | 6 6 6 6 6 |

8 쇰 8 요 ð 원 Š g ð g à g

Š

Jan 28 12:23

7

| AKRPLERVFGFIEKSDLGCGAGGGLANANVFHLAGLTFLTNANADDSQENDEPC-KEILR | |
|--|--|
| <pre>bbB Irrswqimerrmdkagqytdkgirkccedgmrdipmryscqrrariitqgencikaiidc 121 :: :: : :: : </pre> | Db 158/ nikcrnalkidkgkkylmwgiss-di-wgekpnisyligkdtwvehwpeaeed : |
| 728 cnhitkireqhridhviglarseleediipeediisrshipgswlwtieelkepekngis 787 1 : : : : : : : : : | Db 1645 qceelgafesmvvygc 1661 : : : Qy 1660 FIANLDEFAEDIFINCC 1676 |
| 788 tkvmníflkdsíttweilavsladkkgicvadpyeirvmqdffidlrlpysvvrneqvei 847 : : : :::: : : 784 -KQL@FALPDSLTTWEIQGIGISNT-GICVADTVKAKVFKDVFIEMNIPYSVVRGEQIQL 841 | RESULT 7 ID CO3 CAVPO STANDARD; PRT; 1666 AA. |
| 848 ravlfnyreqe-elkvr-ve-l-l-hnpafcsmataknryfqtikippkssvavpyvi 900 :::: :: : : : : :: 842 KGTVYNYRISGMQFCVKMSAVEGICTSESPVIDHQGTKSSKCVRQ-KVEGSSSHLVTFTV 900 | |
| 901 vplkigqqevevkaavfnhfisdgvktlkvvpegmrinktvaihtldpeklgqggv-qk 959 | |
| 960 vdvp-aadisdqvpdtdsetrii-1qgspvvqmaedavdgerlkhlivtpagcgeqmmig 1017 : | |
| 1018 mtptviavhyldqteqwekfgiek-rq-ealel-ikkgytqqlafkqpssayaafnnrpp 1074 | RA AUERBACH H.S., BUNGER R., DODDS A., COLTEN H.R.; RL J. CLIN. INVEST. 86:96-106(1990). RN [2] RP SEQUENCE OF 676-753. |
| 1075 stwltayvvkvfslaanliaidshvlcgavkwlilekqkpdgvfqedgpvihqemiggfr 1134 : :: :: : : : :: : | |
| 135 -nakeadvsltafvlialqeardicegqvnslpgsinkageyieasymnlqrpytvaiag 193 | o z i i m |
| 1194 yalalmnkleepylgkfint-ak-drnr-w-eep-dgql-ynveats 1234 : : : : : : : : : : | 1. |
| 1235 yallalllikdfdsvppvvrwlnegryygggygstqatfmvfqalagyqtdvpdhkdlnm 1294 : | CC +!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVA CC -!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVA CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY CC BOND, C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA |
| 1295 dvsfhlpsrasattfrllwengnllrseetkqneafsltak-gkgrgtlsvvavyhaklk 1353 : ::: :: :: : :: | CC RELEASING C3A ANAPHYLATOXIN 6 GENERATING C3B (BETA CHA CC CHAIN). CC -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN. CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMP |
| 1354 skvtckkfdlrvsirpapetakkpeeakntmfleictky-lg-dvdat-msildismm 1408 | |
| 1409 tgfapdtkdlellasgvdryiskyemnkafsnkntlijylekishteedcltfkvhqyfn 1468 :::: ::: :: :: : : : : : : | CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. DR EMBL; M34054; G305335; DR PIR; A37156; A37156. DR PIR; S03375; S03375. |
| 1469 vgliqpgsvkvysyynleesctrfyhpekddgmlsklchsemcrcaeencf-mqqsqe-k 1526 :: :: | |
| 1527 inlnvrldkacepgvdyvykteltnikllddfdeytmtiqqviksgsdevqagqqrkfis 1586 | KW INFLAMMATORY RESPONSE; GLYCOPROTEIN; SIGNAL. FT SIGNAL |

유

Š

a

õ 요 ð નુ õ

Š

셤

g

ð

유

\$

쇰

ð

g

Š

THOMAS M. L., TACK B.F.;
BIOCHEMISTRY 22:942-947(1983).

-!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.

REACTION IN BOTH CLASSICAL AND ALTERNATIVE, VIA ITS REACTIVE THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES.

-!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN, RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA' 603 KYTCTNA-ELVKGRQYLIMGKEALQIKYNFSFRYIYPLDSLTWIEYWPRDTTC-S-SCQA 1659 587 hikcrnalklqkgkkylmwglss-dl-wgekpntsyiigkdtwvehwpeaeecqdqkyqk 1644 !- SIMILARITY: TO C4, C5 AND ALPHA-2-FRACKOGLAGORITHM.
!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C3,
C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND UKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; .!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. MBL; M34054; G305335; -. OMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN). ERBACH H.S., BURGER R., DODDS A., COLTEN H.R.; CLIN. INVEST. 86:96-106(1990). 1-0CT-1989 (REL. 12, CREATED) 1-UUN-1994 (REL. 29, LAST SEQUENCE UPDATE) 1-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) PRT; 1666 AA EDLINE; 89113342. ERARD N.P., LIVELY M.O., GERARD C.; ROTEIN SEQ. DATA ANAL. 1:473-478(1988) AVIA PORCELLUS (GUINEA PIG) 660 FLANLDEFAEDIFLNGC 1676 645 qceelgaftesmvvygc 1661 BASOPHILIC LEUKOCYTES STANDARD; EQUENCE OF 993-1032. .. <u>=</u> EQUENCE OF 676-753. UTHERIA; RODENTIA. EQUENCE FROM N.A. EDLINE; 83178889. 90307998 O3 CAVPO MEDLINE;

US-08-487-283A-2.rsp

US-08-487-283A-2.rsp

Jan 28 12:23

Jan 28 12:23

| 1666 COMPLEMENT 33. CASA MARTIATORY 13. CASA MARTIATORY 13. CASA MARTIATORY 14. CASA MARTIATORY 14. CASA MARTIATORY 15. CASA MARTIATORY 15 | DOME_LEMENT C3. | | 483 akir | 495 -KIT | 543 evva | 553 ELVS | 603 kgyf | : : 607 SAVY | 663 gldc | PEPC | 723 eacv | 722 PRCI | 783 İkep | 779 L-VP | | | 897 ssva 892 SSHL | 957 lyqg | : 949 I-YG | 1014 gcge | 1006 GSAE | 1071 yaaf : | 1066 YSVW | 1131 hqem : 1126 PIKL | 1190 rpyt | 1184 STFT | 1235 1244 TARM | 1291 vpdh : | 1304 VKQL | 1350 v.avy |
|--|--|---|----------|--|-------------|--------------|------------|-------------------|------------|--------------------------|------------|------------|----------|------------|--|-------------|---------------------------------------|--------------|--------------------------|--------------|----------------------------|----------------------|--------------|--|--------------|--------------|---|------------------------------|--------------|--|
| 166 COMPLEMENT C3. 167 BETA CHAIN. 166 CALPHA CHAIN. 166 CALPHA CHAIN. 166 CALPHA CHAIN. 166 CALPARE (BY C3 CONCERTASE). 167 CLEANAEE (BY C3 CONCERTASE). 168 SIMILARITY. 173 ANAPHYLATOXIN-LIKE. 173 ANAPHYLATOXIN-LIKE. 174 SIMILARITY. 167 BY SIMILARITY. 168 SIMILARITY. 169 SIMILARITY. 160 BY SIMILARITY. 151 BY SIMILARITY. 151 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 166 BY SIMILARITY. 167 BY SIMILARITY. 168 BY SIMILARITY. 168 BY SIMILARITY. 164 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 165 BY SIMILARITY. 166 BY SIMILARITY. 166 BY SIMILARITY. 166 BY SIMILARITY. 167 BY SIMILARITY. 166 BY SIMILARITY. 167 BY SIMILARITY. | 1666 | | qo | δy | qq | Qy | qq | β | QD | Qy | QQ | Qγ | qa | δλ | qq | À. | g & | qu | λΌ | | | | | | | | | | | |
| 166 COMPLEMENT C3. 671 BETA CHAIN. 753 C3A NARPHYLATOXIN. 754 C3A NARPHYLATOXIN. 755 C3A NARPHYLATOXIN. 754 C1EANACE (BY C3 CONCERTASE). 8212 INTERCHAIN (BY SIMILARITY). 822 SIMILARITY. 733 ANAPHYLATOXIN-LIKE. 734 ANAPHYLATOXIN-LIKE. 735 BY SIMILARITY. 737 BY SIMILARITY. 737 BY SIMILARITY. 738 BY SIMILARITY. 739 BY SIMILARITY. 730 BY SIMILARITY. 731 BY SIMILARITY. 732 BY SIMILARITY. 733 BY SIMILARITY. 734 BY SIMILARITY. 735 BY SIMILARITY. 736 BY SIMILARITY. 737 BY SIMILARITY. 738 BY SIMILARITY. 739 BY SIMILARITY. 730 BY SIMILARITY. 731 BY SIMILARITY. 732 BY SIMILARITY. 733 BY SIMILARITY. 734 BY SIMILARITY. 735 BY SIMILARITY. 736 BY SIMILARITY. 737 BY SIMILARITY. 738 BY SIMILARITY. 746 BY SIMILARITY. 746 BY SIMILARITY. 746 BY SIMILARITY. 747 BY SIMILARITY. 748 BY SIMILARITY. 740 BY SIMILARITY. 740 BY SIMILARITY. 741 BY SIMILARITY. 741 BY SIMILARITY. 742 BY SIMILARITY. 742 BY SIMILARITY. 744 BY SIMILARITY. 745 BY SIMILARITY. 745 BY SIMILARITY. 746 BY SIMILARITY. 741 BY SIMILARITY. 742 BY SIMILARITY. 744 BY SIMILARITY. 745 BY SIMILARITY. 745 BY SIMILARITY. 745 BY SIMILARITY. 746 BY SIMILARITY. 747 BY SIMILARITY. 748 BY SIMILARITY. 748 BY SIMILARITY. 749 BY SIMILARITY. 740 BY SIMILARITY. | 1666 | *************************************** | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| 1666 | 1666 | | | | | | | | | | | | | | | | | 2 | 6 | 41 | 38 | 00 | 96 | 59 55 | 15 | 15 | 75 75 | 22 | 35 | 82 |
| 1666 | 30 1666 COMPLEMENT C3. 30 671 BETA CHAIN. 676 1666 ALPHA CHAIN. 676 1666 ALPHA CHAIN. 754 1666 C3B (ALPHA CHAIN). 754 1666 C3B (ALPHA CHAIN). 754 1666 C3B (ALPHA CHAIN). 755 782 175 C1ENAGE (BY C3 CONVERTASE). 750 666 BY SIMILARITY. 751 753 754 SIMILARITY. 752 753 ANDHYLAPOXIN-LIKE. 753 754 1666 BY SIMILARITY. 754 1666 BY SIMILARITY. 755 753 ANDHYLAPOXIN-LIKE. 755 753 754 SIMILARITY. 756 755 821 SIMILARITY. 757 731 731 BY SIMILARITY. 758 1593 BY SIMILARITY. 759 144 944 944 SIMILARITY. 750 1056 BY SIMILARITY. 751 1015 1018 BY SIMILARITY. 752 1593 BY SIMILARITY. 753 734 1462 BY SIMILARITY. 754 1666 BY SIMILARITY. 755 1593 BY SIMILARITY. 756 1669 BY SIMILARITY. 757 101 1560 1669 BY SIMILARITY. 758 1510 BY SIMILARITY. 758 1510 BY SIMILARITY. 759 101 1640 1649 BY SIMILARITY. 750 1040 1664 BY SIMILARITY. 751 101 150 BY SIMILARITY. 752 1593 BY SIMILARITY. 753 101 BY SIMILARITY. 754 1666 BY SIMILARITY. 755 110 101 BY SIMILARITY. 756 1660 BY SIMILARITY. 757 101 1640 1649 BY SIMILARITY. 758 101 1640 1649 BY SIMILARITY. 759 101 1640 1649 BY SIMILARITY. 750 101 1031 BY SIMILARITY. 751 101 1640 1649 BY SIMILARITY. 752 1593 BY SIMILARITY. 753 101 101 BY SIMILARITY. 754 101 1640 1649 BY SIMILARITY. 755 101 1640 1649 BY SIMILARITY. 757 101 1640 1649 BY SIMILARITY. 758 101 1640 1649 BY SIMILARITY. 759 101 1640 1649 BY SIMILARITY. 750 102 1620 POTENTIAL. 750 102 1031 BY SIMILARITY. 751 101 101 101 101 101 101 101 101 101 1 | | | | | | | | | | | | | | | | | tvitsatgy 8 | | tdktiytpg 1 | TOKPVYTPD 1: | wnipelvnm 2 | FKIPSNPRY 1 | | ldgvqpsrp 3 | | | | | |
| 1666 | 30 1666 COMPLEMENT C3. 50 676 1666 ALPHA CHAIN. 676 1666 ALPHA CHAIN. 773 754 1666 C3B (ALPHA CHAIN. 774 1666 C3B (ALPHA CHAIN. 775 821 CLEAVAGE (BY CS) 10 630 666 BY SIMILARITY. 10 630 666 BY SIMILARITY. 10 630 666 BY SIMILARITY. 10 10 630 732 BY SIMILARITY. 10 10 639 732 BY SIMILARITY. 10 10 10 10 10 10 BY SIMILARITY. 10 10 10 10 10 BY SIMILARITY. 10 10 10 10 10 BY SIMILARITY. 10 10 10 10 BY SIMILARITY. 10 10 10 10 BY SIMILARITY. 11 10 10 10 BY SIMILARITY. 12 10 10 10 BY SIMILARITY. 13 10 10 BY SIMILARITY. 14 10 10 10 BY SIMILARITY. 15 10 10 BY SIMILARITY. 16 10 10 BY SIMILARITY. 17 10 10 10 BY SIMILARITY. 18 11 10 10 BY SIMILARITY. 19 140 BY SIMILARITY. 10 1540 1649 BY SIMILARITY. 10 1540 1649 BY SIMILARITY. 10 1540 1649 BY SIMILARITY. 11 10 10 10 BY SIMILARITY. 12 10 10 BY SIMILARITY. 13 10 10 BY SIMILARITY. 14 10 10 BY SIMILARITY. 15 10 10 BY SIMILARITY. 16 166 AA; 186487 MH; D8403BF CR 17 10 10 BY SIMILARITY. 18 11 | | | IN. | CONVERTASE) | SIMILARIII). | ive: | | | | | | | 2). | . 3). . 3). | 532; | | fpakknvlssek | :1 11 11 YPDKKFSYSSGH | lvslqsgylfiq | ::::::!!!! PITYDNGFLFIH | snnqhgilple- | EIDHIGIISFPD | kfy-yiddpkgl : : : WFIGY-KNFKNF | gsgevvlærqvl | SIAQVTFDSETA | nftktpkyfkpa :: :: : NLVATPLFLKPG | -gvarlsintpn : : : | OGVASFVLALPS | elkpgetinvnf |
| 0 1666 COWP 6 1666 ALPH 6 1666 ALPH 73 6 1666 C3B 753 C3A 754 1666 C3B 755 C13B 755 C13B 755 C15B 757 B1 1NTE 8 1517 BY 5 733 BY 5 734 1462 BY 5 735 BY 5 735 BY 5 735 BY 5 737 BY 5 737 BY 5 737 BY 5 738 BY 5 738 BY 5 738 BY 5 739 BY 5 731 BY 5 731 BY 5 731 BY 5 732 BY 5 733 BY 5 734 1462 BY 5 735 BY 5 737 BY 5 737 BY 5 738 BY 5 738 BY 5 739 BY 5 731 BY 5 732 BY 5 733 BY 5 733 BY 5 734 1462 BY 5 735 BY 5 735 BY 5 737 BY 5 737 BY 5 74 1462 BY 5 74 1163 BY 5 74 1462 BY 5 74 1163 BY 5 74 1164 BY | 30 1666 COMP 30 671 BETA 676 1666 ALPH 676 1666 ALPH 676 1666 C3B 754 1666 C3B 754 1666 C3B 755 821 INTE 10 630 666 BY S 10 657 821 INTE 10 699 732 BY S 10 160 1649 BY S 10 1640 1649 BY S 10 1520 1630 BY S 10 1540 1649 BY S 11 151 11 11 11 11 BY 11 151 11 11 11 11 BY 11 11 11 11 11 BY 11 11 11 11 11 BY 11 11 11 11 INTE 11 11 11 INTE 11 11 INTE 11 INT | | CHAIN. | A CHAIN. ANAPHYLATOX (ALPHA' CHA | VAGE (BY C3 | IMILARITY. | IMILARITY. | IMILARITY. | IMILARITY. | IMILARITY. IMILARITY. | IMILARITY. | IMILARITY. | NTIAL. | N (IN REF. | ING (IN REF. ING (IN REF. | D8403B3F CR | 2302; DB No. 0.00e+1 Mismatches | gdipvtvtvhd | : ::: EAFDATISIKS | afggtqlekvv. | : VSKHFSKSKRM | dgipikrdtls: | EGSEVDMVE | psfevlvepte : PHFSVSIEPEY | agsltrvvied | QTAMONTMLIN | gipivtapyqil : GIKYVLSPYKL | vqsltqad :: : : | LDPSKSVTRVDI | nnylhlsmprte |
| 100 | 30 1666 6 6 753 75 4 1666 75 75 4 1666 75 75 4 1666 75 75 4 1666 75 75 821 75 6 1666 75 75 821 75 6 1669 75 821 75 105 609 77 101 71 71 73 71 105 609 72 101 1101 71 101 1640 72 101 101 73 101 74 193 75 1015 70 1640 | | COMP | 38.5 | CLEA | BY S | BY S | | | | | | POTE | - Q | | Σi ~ | | tvvleahevq | : : NIVIQVYGYT | gr-klyvyga | NPVSYVYLEV | rtiivtietp :: : | RETVLTFIDP | aefevkeyvl AYFEVKEYVL | dgdqrisl | рікорокемм | gsdmveaers : : GGFSEEAEIP | -v-vtqg-sn : : | TIDVNQETSD | 423 qtkkggipdarqaintmgalpyttmynsnnylhlsmprtelkpgetinvnfhlrsdpnge 482 |
| | 1 1 1 1 1 1 1 1 1 1 | | | | | | | • | | | → • | → • | | | 1 1 1 1 1 1 1 1 1 1 | | i. | nilrlenee | : : : KIFRVGASE | askefksdk | : :: : P-KQLPGG | vdsdllpvg :: | LNDDLKPAK | enspkqvís : KEDFSTTGT | afvifgv-ç | VYITFGIRE | wertvilhs :: : [AVTVIEST | s-pap-hvr | SVPVI LNAC | arqaintmq |

Que Bes Mat

05 05 05 05 05 05 3 B 3 B 3 B 3 B 3 B B 3

δ

ggv-qreeip-aadlsdqvpdtdsetkil-lqgtpvaqmaedavdaerlkhliitps 1013 GTISRRKEFPYRIPL-DLVPKTEI-KRILSVKGLLVGEILSAVLSQEGINILTHLPK 1005 eqnmigmtptviavhyldqteqwekf-g--lekrqealnlinrgytqqlafkqpnwa 1070 yyaktkrkvvcknfdlrvtlkpapdtvkkpqeakstmilgictry-lg-dqdat-ms 1406 Íknrasstwltayvvkvfslaanligidsevlcgavkwlilekqkpdgvfqedgpvi 1130 -veatsyallalllkdfdavppvvrwlneqryygrgygstqatfmvfqalaqyqtd 1290 1361 : || :| |: :: : : |: |: HISTALD TEASHYRGYGNSDYKRIVACASYKPSREESSSGSS 1420 EAELMSVVPVFYVFHYLETGNHWNIFHSDPLIEKQKLKKKLKEGMLSIMSYRNADYS 1065 tlaiagyalaller----lng--atlqk-fl-naateknr-weea-rqk---l-ys- 1234 602 fvlnkkhkltgskiwdvvekadigctpgsgkdyagvftdaglsfksskaglgtagre 662 722 rneqveiravlynyrea--q-slkvr-vellhnpafcslatakkrhtqtv--tigpk 896 avpyvlvplkiglqevevkaavynyfisdqvkktlkvvpeqmrvnktvairtlnpeq 956 | : ::|| || ||:::: : : | : : ||:|| ||: : : ||:| |
LUTFTVIPLEIGLHNINF--SLETWFGKEILVKTLRVVPEGVKRESYSGV-TLDPRG 948 552 665 721 782 778 842 891 ryytylimnkgkllkvgrqprepgqalvvlpmpitkelipsfrlvayytligasaqr adsvwadvrdscvgtlvvkggsgkdgqdkrqqhlprqqmtlriegnqgarvglvavd| ||::::::|||||: | :|: ||: : : : : | : |
RGEQIQIKGTVYNYRTSGMQFCVKMSAVEGICTSESPVIDHQGTKSSKCVRQKVEGS vkafldcctymaqlrqqhrreqnlglarsdmdediipeediisrsqfpeswlwtiee $perngist kt \verb|mniflkdsittweilavslsdkkgicvadp| fevtvmqdffidlrlpy|$: |:|:: :: ::: ::: :| |: |: ::: | | :|: |
LR-LSMDIDVSYKHKGALANYKMT-DRNFIGRPVEVLIANDDLIVSTGFGSGLATVHV :: ----::

쇰

ð

임

ç

ВÞ

à

원 3

ð

US-08-487-283A-2.rsp

| CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND CC BASOPHILIC LEDWOCYTES. CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. DR EMBL; X52477; G56954; DR EMBL; M29866; E31168; ALT_SEQ. DR PIR; A01260, A01260. DR PIR; S15764; 515764. | PROSTIE; PSOUGT7; ALPHA 2 MAC COMPLEMENT PATHWAY; COMPLEMEN INFLAMMATORY RESPONSE; GLYCOP SIGNAL 1 24 CHAIN 25 1663 CHAIN 25 666 CHAIN 671 1663 | PEPTIDE 671 748 C3A C3A CHAIN 749 1663 C3B SITE 748 749 CLE DOMAIN 693 728 ANA DISULFID 626 661 BY DISULFID 694 727 BY DISULFID 694 727 BY DISULFID 707 728 BY | FT DISULFID 873 1513 BY SIMILARITY. FT DISULFID 1101 1158 BY SIMILARITY. FT DISULFID 1389 1489 BY SIMILARITY. FT DISULFID 1389 1489 BY SIMILARITY. FT DISULFID 1506 1511 BY SIMILARITY. FT DISULFID 1518 1590 BY SIMILARITY. FT THOLEST 1010 1013 BY SIMILARITY. FT THOLEST 1010 1013 PROBABLE. FT CARBOHYD 939 939 PROBABLE. FT CARBOHYD 1617 1617 PROBABLE. FT CONFLICT 721 122 IK -> KL (IN REF. 2). SQ SEQUENCE 1663 AA, 186460 MM; 0428CF63 CRC32; | Query Match 18.7%; Score 2255; DB 2; Length 1663; Best Local Similarity 28.7%; Pred. No. 0.00e+00; Matches 478; Conservative 422; Mismatches 665; Indels 102; Gaps 76 Db 28 ysiitpnvirleseetfileahdaggdvpvtvtvqdfl-kkqvltsektvltgatghlnr 86 | Db 87 vfikipaskefnadkgh-kyvtvvanfgatuvekavlusfqsgylfiqtdktiytpgstv 145 :: : : : | 142 KVRVYSIADDIKPAKRETVIJFIDPEGSEVDMVEEIDHIGIISFDDFKIPSNPRYCMM 205 kirafyehapkqtfsaefevkeyvlpsfevlveptekfy-yihqpkqlevsitarflygk | Db |
|--|---|--|---|---|---|--|---|
| 1407ildismmtgfipdtddlkllatgvdryiskyemnkdfskntliiyldkvshseeecls 1464 :: : : : ::: : : : : : | 1524 iq-lpekitlderlekacepgvdyvyktkllkmelsddfdeyimtieqviksgsdevqa 1581 1 | o 1640 cqdeenqqqcqdlgtftenmvvfgc 1664 | 21-JUL-1986 (REL. 01, CREATED) 01-AUG-1990 (REL. 15, LAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) 00MPLEMENT G3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN). C3. RATTUS NORVEGICUS (RAT). EDIRARNOTA, METAZOA; CHORDATA; VENTEBRATA; TETRAPODA; MAMMALIA; EUTHERIA; RODENTIA. [1] SEQUENCE FROM N.A. STRAIN=WISTAR; TISSUE=LIVER; MEDLINE; 9024567; UTSUIA V. COURA M. TERBARA V. | NUCLEIC ACIDS RES. 18:2178-2178(1990). (2) SEQUENCE OF 671-748. MEDLINE; 79062262. JACOBS J.W., RUBIN J.S., HUGLI T.E., BOCARDT R.A., MARIZ I.K., DANIELS J.S., DAUGHADAY W.H., BRADSHAW R.A.; | SEQUENCE OF 1316-1595 FROM N.A. MEDLINE; 89380332. SUNDSTROM S.A., KOMM B.S., PONCE-DE-LEON H., YI Z., TEUSCHER C., LYTILE C.R.; J. B101. CHEM. 264:16941-16947 (1989). | -!- FUNCTION: G3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. ITS PROCESSING BY G3 CONVERTASE IS THE CENTRAL REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS. AFTER ACTIVATION G3B CAN BIND COVALENTLY, VIA ITS REACTIVE. THIOLESTER, TO CELL SURRACE CARBOHURATES OR IMMUNE AGEREATES. -!- SUBUNIT: G3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE | BAND. C3 CONVEXTABE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN, RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA' CHAIN)!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN!- FUNCTION: DERIVED FROM PROTEOLYTIC DECRADATION OF COMPLEMENT C3, C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT |

76;

| ********* | |
|---|-------|
| | |
| | |
| | 3333 |
| 2000000000 | |
| | |
| ********* | |
| ********** | ***** |
| ******** | |
| ******** | |
| ******* | |
| ******** | |
| | |
| 20000000 | ***** |
| | |
| ********* | **** |
| 200000000000000000000000000000000000000 | |
| | |
| ******* | |
| ********* | |
| ******** | |
| | |
| | |
| ********* | **** |
| | |
| | |
| | |
| | |
| | |
| 2000 | **** |
| 10.000 Contract | |
| 100000000000000000000000000000000000000 | |
| | |
| 444444 | **** |
| | |
| 87-283A-2 | **** |
| | |
| | |
| | |
| 333000000000000000000000000000000000000 | |
| | |
| 2.00 | |
| 8 | |
| | |
| | |
| 200000000000000000000000000000000000000 | |
| | |
| | |
| | |
| 333333.00 | |
| *************************************** | |
| | |
| | |
| | |
| | |
| | ₩ |
| ∞ | |
| | |
| | |
| , 3 | |
| | |
| = | k |
| 9 | |
| Ð | |
| 5 | |
| Ð | |
| J | |
| Ð | |
| Ð | |
| n | |
| n | |
| n | |
| fl | |
| n | |
| fl | |
| n | |
| n | |
| A | |
| n | |
| n | |
| n | |
| n | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| Tan 28 12:23 U | |
| | |
| | |

17

| Š | 319 | NNKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLNLVATPLFLKPGIPYPIKVQVKDS | 378 |
|--------|------|--|------|
| q Q | 380 | -dgs-par-rvp-v-vtgg-sdaqaltq-ddgvaklsvntpnnrgpltitvstk | 426 |
| Ś | 379 | PVILNAQTIDVNQETSDLDPSKSVTRV | 438 |
| QC | 427 | | 486 |
| Š | 439 | APDI.PEENQAREGYRAIAYSSI.SQSYLYIDWTDNHKA.IV | 496 |
| q | 487 | | 546 |
| ò | 497 | THYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPSSRLLVYYIVTGEQTA-ELV | 555 |
| qq | 547 | adsvwvdvkdscvgtlvvkgdprdnrqpapghqttlriegnggarvglvavdkgvfvlnk | 909 |
| Š | 556 | | 614 |
| qq | 607 | knkltgskiwdvvekadigctpgsgknyagvfmdagltfktngglqtdqredpecakpaa | 999 |
| ò | 615 | | 673 |
| q | 199 | rrrrsvqlmerrmdkagqytdkglrkccedgmrdipmpyscqrrarlitgges | 726 |
| ò | 674 | IEEIAAKYKHSVVKKCCYDGAC-VNNDETCEQRAAF | 730 |
| පු | 727 | ccnyitklreqhrrdhvlglarsdvdediipeediisrshfpeswlwtieelkepekngi | 786 |
| ò | 731 | CCVVASQLRANISHKD-MQLGRLHMK-TLLPVSKPEIRSYFPESWLWEVH-L-VP-RR | 783 |
| QQ | 787 | | 846 |
| δ | 784 | KQLQFALDDSLTTWEIQGIGISNT-GICVADTVKAKVFKDVFLEMNIPYSVVRGEQIQ | 840 |
| g | 847 | iravlfnyreqeklkvr-vellhnpafcsmatakkryyqti-e-ippkssvavpyvi | 006 |
| à | 841 | RTSGMQFC | 006 |
| a | 901 | vplkiglqevevkaavfnhfisdgvkkilkvypegmrunktvavrtldpehlngggvqre | 096 |
| Š | 901 | LPLEIGLHNINFSLETWFGKEILVKTLRVVPEGVKRESYSGV-TLDPRGIY-GTISRR | 926 |
| g | 961 | dvnaadls-dqvpdtdsetril-lqgtpvaqmaedavdgerlkhlivtpsgcgeqnmigm | 1018 |
| Š | 957 | I-KRILSVKGLIVGEILSAVLSQEGINILTHLPKGSAEA | 1015 |
| qq | 1019 | tptviavhylddtegwekf-glekrgealelikkgytggl | 1075 |
| à | 1016 | VPVFYVFHYLETGNHWNIFHSDPLIEKQKIKKKKIKEGMLSIMSYRNADYSYSVWKGGSAS | 1075 |
| g | 1076 | twltamwsrsfslaanliaidsqvlcgavkwlilekqkpdgvfqedgpvihqemiggfr- | 1134 |
| δ | 1076 | TWLTAFALRVLGQVNKYVEQNQNSICNSLLMLVENYQLDNGSFKENSQYQPIK | 1135 |
| 엄 | 1135 | | 1194 |
| δy | 1136 | EARENSLYLTAFTVIGIRKAFDICP-LVK-IDTALIKADNFLLENTLPAGSTFTLAISAY | 1193 |
| g | 1195 | alalmnkleepyltkflnt-ak-drnr-weepqqql-ynweatsy | 1235 |
| ò | 1194 | ALS LGDKTHPQFRS IVSALKREALVKGNPP IYRFWKDNLQHKDSSVPNTGTARMVETTAY | 1253 |
| В | 1236 | allalllikdfdsvppvvrwlnderyyggygstqatfmvfqalaqyradvpdhkdlnmd : : :: :: : : ::: :: | 1295 |

Jan 28 12:23 US-08-487:283A-2.rsp 28

| Qy 1254 ALLTSIANKDINYNPVIKWISEEQRYGGGFYSTQDTINAIEGITEYSLLVKQLR-LSMD 1312 | Db 1296 vslhipsrssptvfrilwesgallrseetkqnegfaltak-gkgggtisvvtvyhakvkg 1354 :: ::: ::: : : :: : : :: : : :: : :: : :: : :: : :: : : | 1355 kttckkfdlrvtikpapetakkpqdakssmildictry-lg-dvdat-msildismmt | Qy 1312 EEVCS-FYLKIDIQDIEASHYRGYGNSDYKRIVACASYKPSREESSSGSSHAVMDISLPT 1430 | Db 1410 gfipdtndlellssgvdryiskyemdkafsnkntliiylekishseedclsfkvhqffnv 1469 | Qy 1431 GISANEEDIKALVEGVDQLFTDYQI-KD-GHVILQLNSIPSSDFLCVRFRIFELFEV 1485 | Db 1470 gliqpgsvkvysyynleesctrfyhpekddgmlsklchnemcrcaeencf-mhqsqd-qv 1527 | PATETVYEYHRPDKQCTMFY-S-TSNIKIQKVCEGAACKCVEADCGQ | Db 1528 slnerldkacepgvdyvyktklttielsddfdeyimtieqviksgsdevqagqerrfish 1587 | PEIAYAYKVSITSITVENVFVKYKATLLD | Db 1588 vkcrnalklqkgkgylmwglss-dl-wgekpntsyiigkdtwvehwp 1632 | GKEALQIKYNFSFRYIYPLDSL | 301 | P98094; | DT 01-NOV-1995 (REL.:32, CREATED) DT 01-NOV-1995 (REL. 32, LAST SEQUENCE UPDATE) | DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) DE COMPLEMENT C3 (CONTAINS: C3A ANAPHYLATOXIN) (FRAGMENT). | | | OC AGNATHA (CYCLOSTOMATA). RN [1] | SEQUENCE | | | EMBO J. 11:829-83/(1992)!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF | | CC IMMUNE ACCRECATES. CYCLOSTOMATES C3 APPEARS TO REPRESENT THE | | - ¦·· | | EMBL; 211596; -; NOT A | | | FT NON TER 1 1 COMPLEMENT C3. | <1 633 634 1336 | CHAIN 1343 | 17/ 1CO |
|--|--|---|---|---|--|---|---|---|-------------------------------|--|------------------------|------------|---------|--|--|---|-----|--------------------------------------|----------|------------|------|---|-----|---|-----|------------------|------|------------------------|-----|------------|-------------------------------|--------------------|------------|---------|
| ð | a a | 8 8 | Qy | ΩP | ργ | ΩP | ογ | ପ | δ | ф | Qy | REST 1D | AC. | ro ro | DT DE | S | 8 S | S & | 짪 | X & | RA : | 물 | 8 8 | 38 | 8 5 | 88 | පු ස | i | 5 E | 2 2 | F | : E | : L : | 7 3 |
| | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |

| | • | 8 | æ | |
|-----------|----|----------|-----|----|
| r | | v | | |
| ٠. | 20 | ٠. | 10 | ٠. |
| × | × | | 86 | |
| 0 | × | | 83 | ** |
| × | × | | ø | |
| ۰ | | × | | |
| × | × | ** | 00 | |
| | 90 | 0 | 22 | |
| Ω | ۰ | 9 | | |
| х | 33 | × | 22 | |
| ٠ | ٠ | × | ж | • |
| × | × | × | × | |
| ۰ | | × | × | |
| × | 8 | ** | 88 | * |
| | 93 | | 99 | |
| | 8 | 88 | 60 | |
| | | × | 99 | ٠. |
| ٠ | × | × | 83 | |
| × | 88 | | 00 | × |
| * | 80 | × | × | ** |
| × | • | × | w | × |
| ۰ | м | × | ж | ** |
| × | × | × | ø | × |
| ۰ | ж | × | ж | ** |
| 9 | 82 | × | | |
| × | × | ж | œ | |
| ۰ | 2 | 88 | 88 | |
| × | 8 | ж | 99 | 88 |
| | × | × | × | ** |
| ٠ | × | ۰ | ۰ | 8 |
| ٠ | | | 8 | |
| ۰ | | | ۰ | |
| ۰ | | | ٠ | |
| × | 0 | 8 | 0 | 00 |
| 0 | × | | × | |
| v | 2 | | 88 | |
| × | × | | 99 | |
| ۰ | | | 83 | |
| × | 8 | 88 | o | ** |
| | 8 | œ | w | |
| ۰ | с | 2 | | |
| | Р. | ĸ | | |
| ۰ | н | Ψ. | 43 | ** |
| х | ٠ | | | × |
| •1 | | • | œ | ٠. |
| £ | | ч | 88 | |
| × | | w | æ | ٠. |
| ۰ | 9 | - | | |
| ~ | а | œ | × | • |
| ۰ | 9 | • | 8 | |
| £ | • | | 83 | |
| | ٠. | 4 | × | |
| G | х | 3 | | |
| ö | ٠ | S. | 0.0 | ٠. |
| • | ٠. | | × | ٠. |
| × | х | | | о. |
| ₽ | ٠, | | × | ж. |
| | | | | |
| | ü | × | 8 | |
| Ċ | ü | ۶ | ٠ | |
| ٤ | ÿ | è | i | |
| ١ | ľ | È | | |
| ۱ | į | ì | | |
| ٠ | į | ì | | |
| ç | į | ì | | |
| ۶ | × | è | | |
| ٤ | × | 2 | | |
| 5 | X | Š | | |
| 6 | X | ? | | |
| • | | Š | | |
| • | X | ? | | |
| C | | ? | | |
| C* CC C** | × | } | | |
| C* C* C** | | 2 | | |
| | X | | | |
| | | | | |
| | | } | | |
| | | } | | |
| | | } | | |
| | | | | |
| | | | | |
| | | 3 | | |
| | | 2 | | |
| | | 3 | | |
| | | 3 | | |
| | | 2 | | |
| | | 3 | | |
| 5 C. | | 3 | | |
| | | 2 | | |
| | | 2 | | |
| | | 2 | | |
| | | 2 | | |
| | | 3 | | |
| | | 2 | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | 20 EZ.23 | | |

| -LIKE. | DING. | INTERCHAIN (BY SIMILARITY). | | | | | | | • | | | | | | | CRC32; |
|--------------------|--------------------|-----------------------------|-------------|------------|------------|----------|------------|----------|----------|----------|----------|------------|------------|------------|------------|-----------------|
| ANAPHYLATOXIN-LIKE | PROPERDIN-BINDING. | TERCHAIN (B | SIMILARITY. | SIMILARITY | SIMILARITY | | SIMILARITY | | | - | ٠, | SIMILARITY | SIMILARITY | SIMILARITY | SIMILARITY | C3B9D702 CRC32; |
| æ | id. | ñ | BY | 6 | B | BY | a | BY | BY | BY | BY | BY | æ | æ | £6 | 181747 MM; |
| 694 | 1416 | 780 | 628 | 989 | 693 | 694 | 1474 | 1114 | 1451 | 1420 | 1472 | 1551 | 1616 | 1607 | 973 | Ä |
| 658 | 1406 | 526 | 593 | 658 | 629 | 672 | 836 | 1062 | 1321 | 1356 | 1467 | 1479 | 1499 | 1598 | 970 | 1620 |
| DOMAIN | DOMAIN | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | DISULFID | THIOLEST | SEOUENCE |
| FT | 딦 | F | FI | ΕŢ | F | FT | FT | FI | FI | FT | FT | FT | FT | FI | FI | So |

88; Gaps Query Match 17.5%; Score 2106; DB 2; Length 1620; Best Local Similarity 29.6%; Pred. No. 0.00e+00; Matches 466; Conservative 416; Mismatches 588; Indels 104;

- 93 ipltsrrglvfaqtdqpiytpnndvnirlfpvtrqlnpilsslvvdimnpdgvvvdriek 152 용
- ð
- g
- Š
- 9 δ
- 265 feqgeamhtl-rqkhile-qypdpklllgqslyveasvissdageiensilddipivasp 322 g
 - : | | |: : : | | | | |292 LINGIAQVIFOSERAEIPGIKYVISP 351 ð 8
- ŏ
- 셤
 - õ
- 436 hrevkspgehivfdvfiksaakdhvlhfnylmisngkihnflgegrkgdtt--svslllt 493 :: | | | | : | | | : | | | : | | | : | | | 472 NHKALLVGEHINIIVTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVT 531 셤
 - õ
- pelvpqfrlvaffilp---sgelvadsiiidvkdschaklsldvaggkrlfsprdnvnfd 550 464 요 δ
- 551 lsgesdswavgvvdkaayvldkknkltankvykameasdlgcsvgsgktgplvfrdagl 610 5 õ
- 611 aimakeisgmddvkdpgcpnghtrrkrelvleiaiekasty-paelrkccrdaaiesplr 669 임
 - TFLTNANADDSQENDEPC-KEILRPRRTLQKKIE-EIAAKYKHSVVKKCCYDGACVNN-D 708 652 ð
- 670 laceertkhihdegegcqetflecckhveeelliameeededlgraggedfmigesqvvi 729 셤

:: -= -

US-08-487-283A-2.rsp Jan 28 12.23

8

| ογ | 709 E | ETCEQRAARI-SLGPRCIKAFTECCV-VASQLRANISHKDMQLGRLHMKT-LLPVSKPEI | 765 |
|---------|---------|--|-----|
| QQ | 730 E | shfpesfmweiiklsrsaengksritkkmpdsittwdiqavevsgskgl | 789 |
| δ | 766 R | SYFPESWLWEVHLVPRRK-QLQFALPDSLTTWEIQGIGISNT-GICVA | 819 |
| qq | | alkqyeqvelrvviynymnqdvkgeiqvkcgdqict-daeq :: : :: : : | 846 |
| ý | 820 FI | VVRGEQIQIKGTVYNYRTSGMQFCVKMSAVEGICTS | 879 |
| g | 847 8 | rfav-ek-nsats-f-sfmvvplsssdssvs-vlarvfgsdvh | 901 |
| δλ | 880 3 | SKCVRQKVEGSSSHLVTFTVLPLEIGLHNINFSLETWFGKEILV-KTLRVVPE | 936 |
| οQ | 902 el | rewsvqpr-rhgg-qqvivvdnetpqnvvpgtemsaflsaqgnlv | 959 |
| Qy | 937 E | GIYGTISRRKEFPYRIPLDLVPRTEIKRILSVKGLLVGEI | 966 |
| ф | 960 i | isnlirlprgcgeqnmmytsitvm-varyinrsdqwnkm-gdpqlkkrsfdf-itsgfas | 101 |
| δy | 997 II | | 105 |
| Dp | 1017 q | aawlhrasstwltafvakvfsqarqlvfipvsei | 107 |
| Qy | 1056 11 | | 111 |
| OP O | 1076 g | ovvhln-mmg-qvtgkvv-ltsfvfiallearesc-inevegftvvve | 112 |
| δ | 1116 6 | YLTAFTVIGIRKAFDICP | 117 |
| qq | 1130 gy | ltsqamngledfplaitayalslw-kv-sdgaakvt-m-h-tl-k-tsgl-qtee | 117 |
| ٥y | 1173 NF | LIENTLPAQSTFTLAISAYALSLGDKTHPQFRSIVSALKREALVKGNP | 123 |
| qq | 1180 il | ihw-gsnkgkaaavestaygllaaiqheegeiaekatnwisgsatfggyfgstgdtvm | 123 |
| ò | 1233 QI | GTARMVETTAYALLT | 129 |
| qq | 1237 a | | 129 |
| δ | 1293 A | EGLTEY-SILVKQLRLSMDIDVSYKHKGALHN-YKMTDK | 134 |
| qq | 1297 at | sgtgggiltf-vkkyrekvvikkd-ckgfsleittnldngvkgrrrgsinpefnvyrfi | 135 |
| δ | 1350 T | FGSGLATVHVTTVVHKTSTSEEVC-SFYLKIDTQ-DIEASHYRGYG-NSDYKRI | 140 |
| Qp | 1355 gc | vvmdislptg | 140 |
| ò | 1404 AC | YRPSREESSSGSSHAVMDISLPTGISANEEDIKALVEGVDQLFTDYQIKDGHV | 146 |
| qq | 1410 le | konkdekncvgfrlnqvfesnlvlpv | 146 |
| ð | 1464 LN | SIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHRDDKQCTMFY-STSNIK | 152 |
| ල | 1470 ni | □ . | 152 |
| ď | 1523 A | EEL | 158 |
| В | 1527 tv | vipmhc | 158 |
| ò | 1583 DI | AVAEKDSEITFIKK | 164 |
| q | 1585 ac | dtwfekfplesvc 1598 | |

...

1641 SLIWIEYWPRDITC 1654

```
REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.
AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE
THOLLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGRECATES.
SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG
RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULIDE
BOND. C3 CONVERTABE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN,
RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA'
                                                                                                                                                                                           FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SIMILARITY)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SIMILARITY
                                                                     COMPLEMENT C3-1 (CONTAINS: C3A ANAPHYLATOXIN) (FRAGMENT).
ONCORHYNCHUS MYKISS (RAINBOM TROUT) (SALMO CAIRDNERI).
EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; PISCES; GNATHOSTOMATA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                  C3C FRACMENT (BY SIMILARITY).
C3D FRACMENT (BY SIMILARITY).
C3G FRACMENT (BY SIMILARITY).
C3D FRACMENT (BY SIMILARITY).
C3F FRACMENT (BY SIMILARITY).
C1FRAVAGE (BY C3 CONVERTASE).
                                                                                                                                                                                                                                                                                                                                                      COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (BY
(BY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      INTERCHAIN (BY SIMILARITY)
                                                                                                                                                                                                                                                                                                                              -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (BY FACTOR I)
                                                                                                                                                                                                                                                                                                                   SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (BY FACTOR I)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (BY FACTOR I)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ANAPHYLATOXIN-LIKE.
                                                                                                                                                                                                                                                                                                                                                                                                                            C3A ANAPHYLATOXIN,
C3B (ALPHA' CHAIN)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PROPERDIN-BINDING
                                              01-FEB-1996 (REL. 33, LAST SEQUENCE UPDATE)
01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
            PRT; 1640 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SIMILARITY.
SIMILARITY.
SIMILARITY.
                                                                                                         OSTEICHTHYES; ACTINOPTERYGII; SALMONIFORMES
                                                                                                                                                                                                                                                                                                                                                                                          COMPLEMENT C3.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SIMILARITY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SIMILARITY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SIMILARITY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SIMILARITY
                                                                                                                                                                    LAMBRIS J.D., LAO Z., PANG J., ALSENZ J.;
J. IMMUNOL. 151:6123-6134 (1993).
                                                                                                                                                                                                                                                                                                                                                                                                                 ALPHA CHAIN.
                                                                                                                                 SEQUENCE FROM N.A., AND PARTIAL SEQUENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        CLEAVAGE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CLEAVAGE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CLEAVAGE
                                                                                                                                                                                                                                                                                                                                                                 INFLAMMATORY RESPONSE; GLYCOPROTEIN.
NON TER 1 1 1
CHAIN 1 1640 COMPLEM
                                                                                                                                                                                                                                                                                                                                         EMBL; L24433; G431608; ALT_INIT.
                                   01-FEB-1996 (REL. 33, CREATED)
            STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                          1640
642
642
1640
1640
1640
931
1238
1238
1295
1295
1296
723
1296
703
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         702
703
488
135
464
486
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                563
638
623
164
                                                                                                                                                         MEDLINE; 94065166.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               668
1420
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      536
603
668
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          669
682
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1481
                                                                                                                                              LISSUE=LIVER
LT 10
CO3 ONCMY
                                                                                                                                                                                                                                                                                                        CHAIN)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DISULFID
DISULFID
DISULFID
DISULFID
DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ) ISULFID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ) I SULF I D
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ) I SULF I D
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CARBOHYD
                                                                                                                                                                                                                                                                                                                                                                                                                                                     PEPTIDE
PEPTIDE
PEPTIDE
PEPTIDE
                                                                                                                                                                                                                                                                                                                                                                                                                            EPTIDE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DOMAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            OMAIN
                                                                                                                                                                                                                                                                                                                                                                                                     CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                         CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                                 CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SITE
SITE
SITE
SITE
```

Gaps 429; Mismatches 657; Indels 133; Length 1640; US-08-487-283A-2.rsp 91 BY SIMILARITY. 182104 MW; F4535BB8 CRC32; Score 2002; DB 2; Pred. No. 0.00e+00; 16.64; Best Local Similarity 28.7%; 491; Conservative 991 1640 AA; THIOLEST SEQUENCE Query Match Matches F S

90;

125 yrvfsmtpglepltreifedqevaknkeiavsveimtpenitifreivnpdkgvksgqfk 184 185 lpdivsfgtwhvvtrfgstpgktfssefevkeyvlpsfevsltpakaff-yvddndltvd 243 elvitvktkdpgdprqqtgggtmkalpyrtstknflhvg-vdsn-elkigdpikidlnlg 463 244 itarylygkevtgtg-yvvfgvi-t-tesekksfpaslqrveikdgkgvaclkkeh-itq 299 310 LSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKINLVATPLFIKPGIPY 369 405 430 VLEFNVKTDAPDLPEENQAREGYRAIAYSSLSQSYLYIDWTDNHKALLVGEHLNIIVTPK 489 609 638 pvssrrrravtisdvitsmasky-hglakeccvdgmrdntmgytcdrragyisdgdvcvq 697 670 KEILRPRR--TLOKKIEEIAAKYKHSVVKKCCYDGACVNNDE-TCEQRAARISLGPRCIK 726 aflvcctemaskkieskqdalllsrseedddddaymrsedivsrsqfpeswmwedtnlpe 757 cpaqnkhcestsvirnnflkdsittwqitaislskthgicvadpfemivlkeffidlklp 817 5 vlsapnllrvgsnenifvesqdhvggplnvkimvknhptqskelasksvvldqannfqam 64 : | : : | : | | : | | : | | 310 PIKVQVKDSIDQIVGCVPVILMAQTIDVNQETSDLDPSKSVTRVDDGVASFVILNLPSGVT -tfpkihdlvkqsifvsv8vltegggemveaekrgiqivtspysilfkrtpkyfkpgmpf dvsvyitnp-dn---s-pai-gve-vev---tpd-h-akgvtran-gfakipIntvasat pttipn-hdltymflsrgglvkvg-rfkrggnalvtlsvpvskellpsfrivayyhv-g-SPYIDKITHYNYLLLSKGKITHFGTREKFSDASYQSINIPUTQNMVPSSRLLVYYLVTGE -aadlvadsvwvdikvscmgslkvtstrpkasyeprrafsltitgdpgakvglvavdkgv QTAELVSDSVWLNIEEKCGNQLQVHLSPDADAYSPGQTVSLNMATGMDSWVALAAVDSAV yvlnskhrltqtkiwdtiekhdtgctagggadnmgvfydaglvfetntakgtgirtdpsc :: :: = 329 579 300 490 550 639 758 464 유 õ ò 8 엄 ð 임 ð a δ В ð g g 셤 g 용 용 \$ 8 ò 원 Š ò à

781

ð

용

818 ysavrneglevkailhnysedpiivrvelmengevcssas----kkg-kyrqev--nmdp 870

8 윱 ð В ð 셤 8 유 8 g ð g Š g ð g D Š g õ 셤 ð 셤 8 셤 ð 쇰 ò a 8

OF NONIDENTICAL CHAINS (ALPHA, BETA, AND GAMMA).
POLYMORPHISM: HUMAN COMPLEMENT COMPONENT C4 IS POLYMORPHIC WITH AT
LEAST TWO LOCI, C4A & C4B. 13 ALLELES OF C4A & 22 ALLELES OF C4B
HAVE BEEN DETECTED. THE ALLELE SHOWN HERE IS C4A4. AND, PRIOR TO SECRETION, IS ENZYMATICALLY CLEAVED TO FORM A TRIMER

SUBUNIT: THIS PROTEIN IS SYNTHESIZED AS A SINGLE-CHAIN PRECURSOR

ANAPHYLATOXIN.

ΒY

CINCTION: C4 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE CLASSICAL PATHWAY OF THE COMPLEMENT SYSTEM. IT IS PROCESSED ACTIVATED C1 WHICH REMOVE FROM THE ALPHA CHAIN THE C4A

ANDERSON M.J., MILNER C.M., COTTON G.H., CAMPBELL R.D.; J. IMMUNOL, 148:2795-2802(1992). -!- FUNCTION: C4 PLAYS A CENTRAL ROLE IN THE ACTIVATION

GROUP OF CARBOHYDRATE ANTIGENS.
POLYMORPHISM: THE C4A ALLELES CARRY THE BLOOD GROUP RODGERS WHILE
THE C4B ALLELES CARRY THE BLOOD GROUP CHIDO.
DISEASE: THE C4A6 ALLOTYPE IS TOTALLY DEFICIENT IN HEMOLYTIC

갂. -;

ANTIGENS WHILE C4B ALLOTYPES REACT MORE RAPIDLY WITH THE HYDROXYL

C4A ALLOTYPES REACT MORE RAPIDLY WITH THE AMINO GROUP OF

ACTIVITY.

FUNCTION: DERIVED FROM PROTEDLYTIC DEGRADATION OF COMPLEMENT C4,
FUNCTION: DERIVED FROM PROTEDLYTIC DEGRADAMATORY PROCESS. IT
INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND

US-08-487-283A-2.rsp Jan 28 12:23

EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; MEDLINE; 87080272. YU C.Y., BELT K.T., GILES C.M., CAMPBELL R.D., PORTER R.R.; EMBO J. 5:2873-2881(1986). COMPLEMENT C4 PRECURSOR (CONTAINS: C4A ANAPHYLATOXIN) MEDLINE; 82150875. HARRISON R.A., THOMAS M.L., TACK B.F.; PROC. NATL. ACAD. SCI. U.S.A. 78:7388-7392(1981). BELT K.T., YU C.Y., CARROLL M.C., PORTER R.R.; IMMUNOGENETICS 21:173-180(1985). LAST SEQUENCE UPDATE)
LAST ANNOTATION UPDATE) OF 1-22 AND 1056-1225 FROM N.A. PORTER R.R.; BELT K.T., CARROLL M.C., PORTER R.R.; CELL 36:907-914(1984). MOON K.E., GORSKI J.P., HUGLI T.E.; J. BIOL. CHEM. 256:8685-8692(1981). [6]
STRUCTURAL BASIS OF POLYMORPHISM. CAMPBELL R.D., GAGNON J., PORT BIOCHEM. J. 199:359-370(1981). [2] SEQUENCE OF 20-1741 FROM N.A. 01-NOV-1990 (REL. 16, 01-OCT-1996 (REL. 34, [5] SEQUENCE OF 990-1037. VARIANT C4A6 ALLOTYPE. [4] SEQUENCE OF 957-1044. HOMO SAPIENS (HUMAN) SEQUENCE OF 680-756. EUTHERIA; PRIMATES MEDLINE; 81264286. MEDLINE; 85156269. MEDLINE; 84156544 82182029 MEDLINE; 92242905. TISSUE=LIVER C4A AND C4B. SEQUENCE MEDLINE;

요

ð 셤 ð

쇰 Š

35

| C BASOPHILIC LEUKOCYTES | OCYTES. | è | 301 FD-SETAVKELSYYSLEDIANKYIYTAVTVIESTOOFSEPAFIDGIKYVISDYKIAIIV 357 |
|---|--|------------|---|
| C -!- C4 IS A MAJOR | C4 IS A MAJOR HISTOCOMPATIBILITY COMPLEX CLASS-III PROTEIN. | Ż. | יייייייייייייייייייייייייייייייייייייי |
| C -!- SIMILARITY: TO | SIMILARITY: TO C3, C3 AND ALFHA-Z-MACROGLOBOLLIN. SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN. | QC C | 3/3 KtKrhlvpgaptligalvremsgspasgipvkvsa-tvsspgsvpeagdigmtd-gsgg 43U |
| EWB | 11; | δy | 358 ATPLEIKPGIPYPIKVQVKDSLDQLVGCVPVILNAQTIDVNQETSDLDPSKSVTRVDDCV 417 |
| EMBL; K02403; EMBL; M14824; | 6443671; ALT_SEQ. 6553210; | qq | 431 vsipiiiputiseldlsvsagsph-paiarltvaapp-sg-gogflsjerpds-rppr 484 |
| | | (| |
| DR PIR; A29177; A29177 | | χ̈ | 410 ASEVENTESOVIVEENVAIDAEDEEENQAREGIRALAISSESQSIEIIDMIDNAREE 477 |
| | | qq | 485 vgdtlniniravgs-gatfshyyymilsrgqiv-fmnrepkrtltsvsvfvdhhlaps 540 |
| | 8; HUMAN. | ď | 478 VGEHLNIIVTPKSPYIDKITHYNYLILSKGKIIHFGTREKFSDASYQSINIPVTQNMVPS 537 |
| OR MIM; 120790; | | 4 | 541 fufunctur hadh-n-man widiranan and and anternament by order |
| | | a a | TXIVATYY-iigui-pvaiisIIVovqaqacegArtetsva-qakqYriigesvAriiIteros |
| OR. PROSITE; PS00477; P | PROSITE; PS00477; ALPHA 2 MACROGLOBULIN. COMPLEMENT PARHWAY: PLASMA: CIXCORBOTEIN: MHC III: SIGNAL: | δλ | 538 SRLLVYYIVTGEQTAELVSDSVMLNIEE-KCGNQLQVHLSPDADAYSPCQTVSLNMATGM 596 |
| | INFLAMMATORY RESPONSE; POLYMORPHISM; DISEASE MUTATION. | qq | 596 lalvalgaldtalyaagskshkplnmgkvfeamnsydlgcgpggddsalqvfqaaglaf- 654 |
| 50 7 | | δ | 597 DSWVALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGGLNNANVFHLAGLTFL 654 |
| 680 1451 | 1443 ALPHA CHAIN. 1741 GAMMA CHAIN. | QQ | 655 sdgdqwtlsrkrlscpkekttrkkrnvnfgkaineklgqyasptakrccqdgvtrlpmmr 714 |
| T PEPTIDE 680 | 756 C4A ANAPHYLATOXIN. 736 ANAPHYLATOXIN-1.1KF. | ò | |
| | | X 1 | |
| T DISULFID 703 | 735 BY SIMILARITY. 736 BY SIMILARITY. | අු | 715 scegraarvgg-pdcrepflsccgfaeslrkksrdkggaglqraleilgeedlideddip 773 |
| 1010 | | ογ | 710 TCEQRAARISLGPRCIKAFTECCVVASQLRA-NISHKDMQLGRLHMKTLLPVSKPE 764 |
| | | ପୁ | etvdrfqiltlwlpdslttweihglslsktkglcvatpvql |
| | 1328 POTENTIAL. 1391 : POTENTIAL. | 'n | |
| T VARIANT 477 | 477 R -> W (IN C4A6). 726 P -> 1. (TN C4A3). | · £ | 834 hihirimaurrfen leirnvinnvidknitusuhusnum alciannniannulunana 893 |
| 1073 | D -> G (IN C4A1, | 3 , | *** : : : : : : : : : : : : : : : : : : : : : : : : |
| | 1125 PCPVID -> LSPVIH (IN C4B). 1176 N -> S (IN C4A1, C4B1, C4B3 AND C4B5). | δλ | 824 FLEMNIPYSVVRGEQIQLKGTVYNYRTSCMQFCVKMSAVEGICTSESPVIDHQGTKSSKC 883 |
| 1201 | (IN C4A6, C4A3, C4 | ପ୍ଧ | 894 arp-vafsvvptaaaav-slkvvarg-sfefpvg-davskvlqiekegaihreelvye 947 |
| 1210 | L -> R (IN C4A1, C4B1, C | δ | LP LEIGLHNINFSLETWFGKEILVKTLRVVPE |
| 727 | D -> N (IN REF. 3). | qq | 948 inpidhrgrileipgnsd-pnmip-dgdfnsyv-rvtasdpldtlgse-ga-lspggvas 1002 |
| T CONFLICT 1013 1 SQ SEQUENCE 1741 AA; | 1013 Q $^{-7}$ E (IN KEF: 4 AND 5). ; 192335 MM; 405A8805 CRC32; | δλ | 941 GVTLDPRGIYGTISRRKEFPYRIPLDLVPKTEIKRILSVKGL-LVGEILSAVLSQEGINI 999 |
| Query Match | 14.3%; Score 1723; DB 2; Length 1741; | qu | 1003 11r1prgcgeqtmiylaptlaasryldkteqwst1ppet-kd-havdligk-gymriqqf 1059 |
| Matches 469; Conser | 4 | ٥y | 1000 LTHIPKGSAEAELMSVVPVFVFFYLETGNHWNIFHSDPLIEKQKLKKKKLKEGMLSIMSY 1059 |
| b 139 ghlflqtdqpiyng | t-dtitvm | QQ | 1060 rkadgsyaawisrdsstwitafvikvislaqeqvggspekiqetsnwilsq-qqadqsfq 1118 |
| y 124 GFLFIHTDKPVYTE | | Qy | 1060 RNADYSYSVWKGGSASTWLTAFALRVLGQVNKYVEQNQNSICNSLLMLVENYQLDNGSFK 1119 |
| b 198 fq-ddfvipdiser | fq-ddfvipdisepgtwkisarfedglesnsstqfevkkyvlpnfevkitpgkpyiltvp 256 | qq | rtialhhglavfqdegaeplkq |
| Y 183 ISFPDFKIPSNPRY | : | δ | 1120 ENSQYQPIKLQGTLPVEARENSLYLTAFTVIGIRKA-F-DI-C-PLV-KIDTALIKAD 1172 |
| b 257 ghldemqldiqary | ghldemqldiqaryiygkpv-qqvayvrfglldedgkktffrglesqtklvngqshis 313 | QQ | 1177 sflgekasagllgahaaaitayalsit-kapvdllgvahnnlmamaqetgdn-lywgsvt 1234 |
| y 242 -NFKNFEITIKARY | -NFKNFEITIKARYFYNKVYTEADVYITFGIREDIKDPOKEMMQTAMQNTMLINGIAQYT 300 | Qy | 1173 NFILENTLPAQ-STFTLAISAYALSLGDKTHPQFRSIV-SALKREALVKGNPPIYRFWKD 1230 |
| b 314 løkaefqdaleklno : : : : | 314 lskaefqdaleklnm-gitdlgglrlyvaaaiiespggemeeaeltswyfvsspfsldls 372 : : : : : : | q | 1235 gsqsnavsptpaprnpsdpmpqapalwiettayallhlllhegkaemadqasawltrqgs 1294 : : : |

쇰 ð 9 ð 셤 ð 음 STANDARD;

CO4 MOUSE

1534

요 ð 쇰 õ 셤 3 셤 ð

ð

[1] SEQUENCE FROM N.A. MEDLINE; 85298264.

[2] SEQUENCE FROM N.A. MEDLINE; 87309760.

STRAIN-B12.WR;

[3] SEQUENCE FROM N.A.

US-08-487-283A-2.rsp Jan 28 12:23

```
ANAPHYLATOXIN.
SUBUNIT: THIS PROTEIN IS SYNTHESIZED AS A SINGLE-CHAIN PRECURSOR
AND, PRIOR TO SECRETION, IS ENZYMATICALLY CLEAVED TO FORM A TRIMER
OF NONIDENTICAL CHAINS (ALPHA, BETA, AND GAMMA).
                                                                                                                                                                                            NONAKA M., TAKAHASHI M., NATSUUME-SAKAI S., NONAKA M., TANAKA S.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NONAKA M., NAKAYAMA K., YEUL Y.D., SHIMIZU A., TAKAHASHI M.;
IMMUNOL. REV. 87:81-99(1985).
-!- FUNCTION: C4 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
CLASSICAL PATHANY OF THE COMPLEMENT SYSTEM. IT IS PROCESSED
ACTIVATED CI WHICH REMOVE FROM THE ALPHA CHAIN THE C4A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         C4 IS A MAJOR HISTOCOMPATIBILITY COMPLEX CLASS-III PROTEIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TOSI M., LEVI-STRAUSS M., DUPONCHEL C., MED T.;
PHILOS. TRANS. R. SOC. LOND., B, BIOL. SCI. 306:389-394(1984).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.
COMPLEMENT PATHWAY; PLASMA; GLYCOPROTEIN; MHC III; SIGNAL;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      -!- SIMILARITY: TO C3, C5 AND ALPHA-2-MACROGLOBULIN.
-!- SIMILARITY: CONTAINS ONE AND HYLATOXIN-LIKE DOMAIN.
EMBL, KO0019; E19386; -.
EMBL, MI1296; G39720; -.
EMBL, MI2970; G199267; -.
EMBL, MI2970; G199267; -.
EMBL, MI2979; G397389; -.
EMBL, MI2979; G397887; -.
EMBL, MI799; G397440; -.
                                                                                                                                                                                                                                                                                                                                                SEQUENCE OF 1105-1449 FROM N.A.
MEDLINE; 85166208.
LEVI-STRAUSS M., TOSI M., STEINMETZ M., KLEIN J., MEO T.;
PROC. NATL. ACAD. SCI. U.S.A. 82:1746-1750(1985).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         OGATA R.T., SHREFFLER D.C., SEPICH D.S., LILLY S.P., PROC. NATL. ACAD. SCI. U.S.A. 80:5061-5065(1983).
NONAKA M., NAKAYAMA K., YEUL Y.D., TAKAHASHI M.;
                                                                                                                                                                                                                                                                            PROC. NATL. ACAD. SCI. U.S.A. 81:6822-6826(1984)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             C4A ANAPHYLATOXIN.
                                                                                                                SEQUENCE OF 651-810 AND 924-1083 FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ALPHA CHAIN.
GAMMA CHAIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         BETA CHAIN.

    BIOL. CHEM. 260:10936-10943(1985).
```

BY

| ***** | |
|---------------|--|
| ********** | |
| | |
| | |
| | |
| | |
| | |
| ******* | |
| ********* | |
| ******** | |
| | |
| | |
| ******* | |
| ******* | |
| | |
| | |
| | |
| | |
| ******** | |
| | |
| | |
| ****** | |
| ******* | |
| | |
| ******** | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| ~ 3 | |
| | |
| | |
| | |
| 24 | |
| ~~ | |
| Y () | |
| | |
| | |
| | |
| 00 | |
| | |
| | |
| | |
| | |
| | |
| | |
| X | |
| A00 (000) | |
| | |
| ****** | |
| ******** | |
| | |
| | |
| ******* | |
| ******** | |
| ******* | |
| ******* | |
| | |
| ******* | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| 65 | |
| SZ. | |
| 83 | |
| 2.23 | |
| 12.23 | |
| 12:23 | |
| 8 12:23 | |
| 28 12:23 | |
| 28 12:23 | |
| n 28 12:23 | |
| an 28 12 23 | |
| Ian 28 i 2:23 | |
| Jan 28 12:23 | |
| Jan 28 12:23 | |

| ANAPHYLATOXIN-LIKE. | SIMILARITY. | SIMILARITY. | LARITY. | LARITY. | AL. | | AL. | (IN REF. 4). | (IN REF. 4). | N | (IN REF. 5). | | Z | Z | (IN REF. 5). | z | Z | z | | | (IN REF. 4). | A6cclFCB CRC32; |
|---------------------|-------------|-------------|----------|----------|------------|----------|------------|--------------|--------------|----------|--------------|----------|----------|----------|--------------|----------|----------|----------|----------|----------|--------------|-----------------|
| ANAPHYI | BY SIM | BY SIM | BY SIM | BY SIM | POTENTIAL. | | POTENTIAL. | F -> Y |) 3 ^ 5 | 3 - O | R - G | 01 -> 1 | P -> R | P -> L | 0 -> E | V A | A -> T | K -> N | G - S | R -> K | A -> V | MW; A60 |
| 734 | 726 | 733 | 734 | 1009 | 224 | 743 | 1387 | 132 | 327 | 570 | 720 | 740 | 838 | 993 | 1043 | 1119 | 1190 | 1324 | 1401 | 1442 | 1453 | AA; 192870 P |
| 700 | 700 | 701 | 714 | 1006 | 224 | 743 | 1387 | 132 | 327 | 570 | 720 | 739 | 838 | 993 | 1043 | 1119 | 1190 | 1324 | 1401 | 1442 | 1453 | 1738 |
| DOMAIN | DISULFID | DISULFID | DISULFID | THIOLEST | CARBOHYD | CARBOHYD | CARBOHYD | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | CONFLICT | SEQUENCE |
| F | Ε | 닲 | F | F | 댠 | ፎ | F | FT | ŗ | FJ | F | FŦ | FT | FT | FŢ | FI | FI | F | F | FT | F | δS |

Gaps 111; Indels 143; Length 1738; 407; Mismatches 628; Score 1605; DB 2; Pred. No. 0.00e+00; 13.3%; 27.6%; 449; Conservative Local Similarity Query Match Matches

음 8

254 240 SPPD-FKIPSNPRYGMFIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEP-E-YNFIGY 184 g õ

snsdeiqldiqaryiygkpv-qgvaytrfalmde--qgkrtflrgletqaklvegrthis 311 255 g

300 241 ð 312 301 g Š 372 359 g Š

482 419 429 셤 ò

gdtfilnlqpvgipaptfshyyymiisrgqimamg-rep-rktv-tsvsvlvdhqlapsf 539 |::::|:::||CEHLANITYTPKSPYIDKITHYVILLSKGKIIHFGTREKFSDASYQSINIPVTQNAVPSS 538 483 셤

479

õ

540 yfvayfyhqg-h-p--vanslliniq&rdcegklqlkvd-gakeyrnadmmklriqtdsk 594 597 SELVYYIVTGEQTAELVSDSVWINIEEK-CGNQLQVH! SPDADAYSPGQTVSIANATGMD 염 õ 654 alvalgavdtalyavggrshkpldmskvfevinsynvgcgpggddalqvfqdaglafsd : |||:|||:|| : ||:|||:|| SWVALAAVDSAVYGVQRCAKKPLE--RVFQFLEKSDLGCGAGGGLNNANVFHLAGLFFLT 595 598 윤 ò

US-08-487-283A-2.rsp Jan 28 12:23

33

1116 1416 944 ldp--1-nnlgrtleipgssdpnivpdgdfsslvrvtasepletmgsegalspggvasll 1000 1001 rlpggcaeqtmiylaptltasnyldrteqwsklspet-kd-havdliqk-gymriqqfrk 1057 1061 1175 lgqkasagllgahaaaitayaltlt-kasedlrnvahnslmamaeetgeh-lywglvlgs 1232 | ::::::::|||||||||| | | :::|:::||| | | ::|: |175 ILENTLPAQ-STFTLAISAYALSLGDKTHPQFRSIV-SALKREALVKGNPPIYR---F-W 1228 1329 1530 1475 1590 1121 1117 cpvihramqggl-vgs-detvaltafvvialhhgldvfqdddakqlknrveasitkansf 1174 1351 1471 qdkvvlrptaprsptepvpqapalwiettayall-hlllregkgkmadkaaswlthqgsf 1291 1411 1372 772 1h1r1pisirrfeqfe1rpv1yny1nddvavsvhvtpveg1clagggmmaqqvtvpagsa 892 832 824 884 rtsfpenwlwrvepvdssklltvwlpdsmttweihgvslskskglcvakptrvrvfrkfh hgafrstqdtvvtldalsaywiashtteekalkvtlssmgrnglkthglhlnnhqvkgle 655 gdrltqtre-dlscpkekksrqkrnvnfqkavseklgqysspdakrccqdgmtklpmkrt 656 NANADDSQENDEPC-KEIL-RPRFIL--QKKIEEIAAKYKHSVVKKCCYDGAC-VNNDET cegraarvpqqa-crepflscckfaedlrrnqtrsqahlarnnhnmlqeedlideddilv 1002 HLPKGSAEAELMSVVPVFVFFFFTETGNHWNIFHSDPLIEKQKLKKGKLKEGMLSIMSYRN eelkfslqstisvkvegnskgtlkilrtynvldmknttcgdlgievkvtgaveyawdane ::| || : | : | | : | : | BK-NF-LGRPVEVLL--ND-D-L-IVST-G-FG----S--GLAT-VHVTTVVHKT-STSE gfgasqevvvglvqpssavlydyyspdhkcsvfyaaptksqllatlcsgdvcqcaegkcp 1472 lglsgmaiaditllsgfhalradlekltslsdryvshfetdgphvllyfdsvpttr-ecv .. •• 714 991 833 885 944 773 825 1233 1229 1330 1412 893 1292 1281 1352 염 셤 셤 셤 Б 셤 g à δ g õ ò ð δ 염 ò g В g g Š ò à Š ò P ò ð g ð g ò

#

US-08-487-283A-2.rsp

| a | 1591 rllrølerrvedkdgyrmrfacyyprveygftvkvlredgraafrlfeskitqvlhfrkd 1650 | Qy 1001 | THIPKGSAEAELMSVVPVFYVFHYLETGNHMNIFHSDPLIEKQKIKKKKKEGMLSIMSYR 106 |
|----------------|---|----------|---|
| δ | :: : :: : : : : : : : : : | Db 124 | qpnsayaaflnrapstwltayvvkvfslavnliaidsqvlcg |
| PP PP | 1651 tmasigqtrnf18rasc-r1rlepnkeylimgmdgetsdnkgdpqylldsntwieemp | Qy 1061 | 1061 NADYSYSVAKGCSASTWLTAFALRVLGQVNKYVEQNQNSICNSLIMLVENYQLDNGSFKE 1120 |
| ठे द | 1590 AVAEKDSEITFIKKVTCTNAELVKGRQYLIMGKEALQIKYNFSFRYIYPLDSLTWIEYWP 1649 | | 184 dapvihqemiggqr-nsvekeraltafvlialqeareiceeqvnslaasinksrdflaan 242 |
| a a | 17.1 avanta 05.1 | Vy 1121 | NOVICE IN UNIO I EVERARENO LI LI RETVICI ERABE DI CETTANO I DI ALIMADNE LIGANI I I 10 |
| Ŝ | 1000 RD11C03 1000 | | 243 YMINGLEDYAVALAGYAWAGGASTTIGTATINA LIBRARKANTHIN-WEEPIGG-T |
| RESULT ID C | ILT 13 COSTANDARD; PRT; 726 AA. | | 290 l-ynveassyallallltdfdsvppvvrwlneqryygggygstqatfmgfqala 343 |
| PAC | 01-0CT-1989 (REL. 12, CREATED) | Qy 1239 | ::: : ::: : ::: : : : : |
| 2 2 2 | 01-001-1989 (REL. 12, LAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) COMDITENTAL CALAIN (FRAGMENT) | Db 344 | 344 qyqtdypdhkdlnmvvsiqlpsrsspvkhrivwdsasllrseetkenqgfsltaq-gkgq 402 |
| 8 8 8 | OCH MENERAL CO MALLIN CHANGE IN THE CONTROL OF THE | Qy 1299 | 1299 EYSLLVKQLR-LSMDIDVSYKHKGALHNYKMT-DKNFLGRPVEVLLNDDLIVSTGFGSGL 1356 |
| 3 S S | CONTROLLANDO CONTROLO (NORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; ETITURETA: I ACCMODULA | Db 403 | gtløvvttyfakvkgkvtckkfdlrvniktapetvkkpqdakstmilghctry-lg-ded 460 |
| 8 8 8 | CONTRACTO COM N A | Qy 1357 | |
| ≵ ≵ ≴ | DEJOENCE FROM N.A. MEDLINE 87006901 N.H., TOMITA M., YAMAMOTO K., MIGITA S., SEKIYA T., | Db 461 | at-msildismmtgfvpdtddlnllstgvdryiskyelnkafsnkntliiyldkishs 517 |
| R. R. | NISHIMURA S.; IMMUNOL. INVEST. 15:365-378(1986). | Qy 1416 | SGVDQLFTDYQI-KD-GH |
| 88 | -!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL | Db 518 | reeclafkvhqyfnvgliqpgavkvysyynleetctqfyhpekedgmlsklchkemcrca 577 : :::: : :: : : : :: :: |
| ខ្លួ | REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS. AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE | Qy 1471 | 1471 DFLCVRFRIFELFEVGFLSPATFTVYEYHRPDKQCTMFY-S-TSNIKIQKVCEGAACKCV 1528 |
| ខ្ល | THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES. -!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG | Db 578 | 578 eencf-mqq-1dekitlndr1dkacepgldyvyktklvqveraddfdeylmvventiksg 635 |
| 88 | RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN, | Qy 1529 | |
| ខខ | RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA' CHAIN). | DP 636 | 636 sdevqagqpapfishikcrdalklkdgkhylmwglssdpvgekpntsyiigkdtwvef 693 : : : : |
| S 8 | -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN. EMBL: M32434; G164863: | Qy 1589 | KEALQIKYNFSFRYIYPLDSI |
| 8 8 8 | | | 694 wpekeecqdeenqkhcedlgafaesmvvfgc 724 : : |
| 2 2 1 | COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA; INFEMMATORY RESPONSE; GLYCOPROTEIN. | Qy 1648 | 1648 WPRDTTC-SS-CQAFIANIDEFAEDIFINGC 1676 |
| 1 1 1 | NOW LEK 1 1 ALPHA CHAIN. PURIOR 71 76 ALPHA CHAIN. | RESULT 1 | 14 crannada. Dor. 1472 aa |
| 1 11 1 | • | | נמזנט |
| : E 8 | 680 680 726 AA; 81844 MW; | DT 01-AP | 01-APR-1990 (REL. 14, LAST SEQUENCE UPDATE) 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE) |
| ő | erv Watch 6.6%: Score 796: DB 2: Length 726: | | ALPHA-1-INHIBITOR III PRECURSOR. RATTUS NORVEGICUS (RAT) |
| * of \$2 | Best Local Similarity 26.2%; Pred. No. 5.45e-148; Matches 197; Conservative 197; Mismatches 305; Indels 52; Gaps 37; | | EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; EUTHERIA; RODENTIA. III |
| a 8 | 9 tldpenlggggv-qkeeipsadisdqvpgtesetkil-lggtpvaqmtedaidgerlkhl 66 : : : : : : | | SEQUENCE FROM N.A., AND PARTIAL SEQUENCE. 15SSUE=LIVER; MEDIIVE: 88153707 |
| 2 23 | 67 ivtosacaeanmiamthtviavhvldhteawdkf-slekraealelikkavtaalafk 123 | | BRACIAK T.A., NORTHEMANN W., HUDSON G.O., SHIELS B.R., GEHRING M.R., FEY G.H.; |
| ļ | | RL J. BI | J. BIOL. CHEM. 263:3999-4012(1988). |

g

186 g1kq1s-fs1saepiqgpyki-vilkqs-gvkeehsftvmefv1prfgvdvkvpnaisvy 242 |: ::| | | | | | : : | | | | | | GIISFPDFKIPSNPRYGMWTIKAKYKEDFSTTGTAYFEVKEYVLPHFSVSIEPEYNFIGY 240

181

ð

අු

õ

US-08-487-283A-2.rsp Jan 28 12:23

₩.

\$

| g | 243 de | deiinvtacatytygkpyp-ghvkisl-chgnptfssetksgckeedsrldnngcstq 298 |
|-----|----------------------|---|
| δ | 241 KN | FKNFEIT |
| qq | 299 ev | evnitefqlke-nylk-mhqaf-hvnatvteegtgsefsgsgriever-trnkf-lfl 351 |
| ò | 299 VT | VTFDSETAVKELSYYSLEDLANKYLY1AVTVIE-STGG-FSEFAEIPGIKYVLSPYKLNL 356 |
| qq | 352 ka | kadshf-rhgipffvkvrlvdikgdpipneqvlikardagytnatttdqhglakfsidtn 410 : : : : : : : : : : : : |
| δ | 357 VA | VATPLFLKPGIPYPIKVQVKDSLDQLVGGVPVIINAQTIDVNQETSDLDPSKSVTRVD-D 415 |
| QQ | 411 gi | gisdyslnikvyhkeesscihssctaerhaeah-htay-av-ys-lsksyiyld-tea-g 464 [::: |
| δλ | 416 GV | ASFVIALPSGVTVLEFNVKTD-APDLPEENQAREGYRAIAYSSLSQSYLYIDWTDNHK 474 |
| QQ | 465 vl | ivfhylvmaqgsilqtgnl |
| ٥y | 475 AL | ALLVGE-H-INIIVTPKSPYIDKITHYNYLILSKGKIIHFGTRE-KFSDASYQ 524 |
| qq | 525 al | aleipvefsmvpvakmliytilp-dgeviadsvkfqve-kclrn-kvhlsfspsqslp 579 |
| δy | 525 51 | .NIPVTQNMVPSSRLLVYYIVTGEQTAELVSDSVWINIEEKCGNQLQVHLS-PDADAYS 583 |
| qq | 580 as | asqt-hmrvtaspqslcglravdqsvl-lqkp-e-a-e-lsps-liydlpg-mqds 627 |
| ٥y | 584 PG | GAK |
| Op | 628 n- | n-f-ias-sndpf-ed-edyclmyqpiarek-dvyryvretglmaftnlkiklpty 677 |
| ٥y | 644 NV | FHLAGLIFLTHNANADDSQENDEPCKEILRPRRTLQKKIEEIAAK-YKHSVVKKCCYDG 702 |
| DP | 678 -ci | cntdydmvplavpavaldsstdrgmyeslpvva-vksplpqepprkdppp-kdpvie 732 |
| Qy | 703 AC | VNNDETCEQRAARISLGPRCIKAFTECCVVASQLRANIS |
| QQ | 733 t- | -irnyfpetwiwdlutunssgutelemtupdtitewkagalcisndtglgissvasfqa 791 |
| δλ | 763 PE | PEIRSYFPESMIME-VHIVPR-RKQLQFALPDSLTTWEIQGIGISN-TGICVADTVKAKV 819 |
| ΟP | 792 fg | <pre>apffveltmpysvirgeaftlkatvlnylptslpmavlleaspdf-ta-vpvenngd 847</pre> |
| οy | 820 FK | |
| QQ | 848 sy | -clgangrhtsswlvtpkslg-nvnf-svs-a-eargspgpcgsevatvpetgrkd |
| δy | 880 55 | |
| qq | 903 vk | ckeht |
| δý | 940 S _Č V | V-TIDPRGIYGTISRRKEFPYRIPIDIV-PKTEIKRILSVKGILVGEILSAVL 992 |
| qq | 963 -kı | -kntqm-liqmpygcgeqmwvlfapniyvlkyln-etq-qltekikska-lgy-lrag 1014 |
| δý | 993 SQI | AEAELMSVVP VFYVFHYLETGNHWNIFHSDPLIEKQKLKKKL |
| ΟP | 1015 yq | yqrelnykhkdgsysafgdhngqqqqntwltafvlksfaqarafifideshitdaftwls 1074 |
| ò | 1053 ML | MLSIMSYRNADYS-YSVWKG-GSASTWLTAFALRVLGQVNKYVEQNQNSICNSLLWIV 1108 |
| qq | 1075 kg | kq-qkdsgcfrssgsllnnamkggvddeitlsa-yit-mallesslp-dtdpvvskalsc 1130 |
| ο̈́ | 1109 EN | ENYQLDNGSFKENSQYQPIKLQGTLPVEARENSLYLTAFTVIGIRKAFDICPLVKIDTAL 1168 |

OVERBERGH L., TORREKENS S., VAN LEUVEN F., VAN DEN BERGHE H.; LAST SEQUENCE UPDATE)
LAST ANNOTATION UPDATE) PRT; 1476 AA. [1] SEQUENCE FROM N.A., AND SEQUENCE OF 28-57. MURINOGLOBULIN 1 PRECURSOR (MUG1) CREATED) STANDARD; 01-DEC-1992 (REL. 24, 01-NOV-1995 (REL. 32, 01-NOV-1995 (REL. 32, MUS MUSCULUS (MOUSE) EUTHERIA; RODENTIA. MEDLINE; 91358495. MUG1 OR MUG-1 TISSUE-LIVER A2MG MOUSE P28665; 15

1. BIOL. CHEM. 266:1603-16910(1991).

-!- FUNCTION: A PROTEINASE ACTIVATES THE INHIBITOR BY SPECIFIC PROTEOLYSIS IN THE BAIT RECION, WHICH, BY AN UNKNOWN MECHANISM LEADS TO REACTION AT THE CYSTEINYL-CLUTAMYL INTERNAL THIOL ESTER SITE AND TO A CONFORMATIONAL CHANGE, WHEREBY THE PROTEINASE IS TRAPPED AND/OR CONALENTLY BOOND TO THE INHIBITOR. WHILE IN THE TETRAMERIC PROTEINASE INHIBITORS STERIC INHIBITION IS SUFFICIENTLY ACTIVATED GLUTAMYL RESIDUE OF THE ORIGINAL THIOL ESTER AND A TERMINAL AMINO GROUP OF A LYSINE OR ANOTHER NUCLEOPHILIC GROUP ON THE PROTEINASE, FOR INHIBITION TO BE EFFECTIVE. PIR; A41185, A41185. PROSITE; PSO0477; ALPHA 2 MACROGLOBULIN. SERINE PROTEASE INHIBITOR; GLYCOPROTEIN; PLASMA; BAIT REGION; SIGNAL; MULTIGENE FAMILY. -!- SIMILARITY: TO OTHER PROTEINS OF THE ALPHA-MACROGLOBULIN FAMILY, INCLUDING COMPLEMENT COMPONENTS C3, C4, AND C5. STRONG, MONOMERIC FORMS NEED A COVALENT LINKAGE BETWEEN THE EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA; MURINOGLOBULIN 1. BAIT REGION (APPROXIMATELY). -!- TISSUE SPECIFICITY: PLASMA -!- SUBUNIT: MONOMER. EMBL; M65736; G199891; -. 27 1476 734 28 677 CHAIN DOMAIN SIGNAL

| ****** |
|---|
| |
| |
| |
| ******** |
| |
| ******* |
| |
| ********* |
| |
| |
| |
| ********* |
| |
| ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, |
| |
| •••••• |
| |
| ••••• |
| ******* |
| ********** |
| ******* |
| |
| |
| •••••• |
| |
| |
| |
| |
| |
| v.: 4.2000 |
| |
| |
| |
| |
| |
| A 40 (000) |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| 99 |
| ఈ |
| ĕ. |
| Ħ |
| Ĭ |
| 3 5 |
| 89 129 |
| <u> </u> |
| S C S S |
| 285 C8-08 |
| SP-SS |
| SP-SIT |
| 89 29 |
| 868J |
| US-08-487-283A-2.rsp |
| 38-83 18-08 |
| 36-SJ |
| US-08 |
| 78-0S |
| 30-SN |
| US-08 |
| 30-SN |
| US-08 |
| 78-98 118-08 |
| OS-OS |
| US-08 |
| SO-SI) |
| 30-SN |
| 30-SN |
| 30-SN |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| Jan 28 12.23 US-08 |

æ

| 48 86 BY SIMILARITY. 269 288 BY SIMILARITY. 269 288 BY SIMILARITY. 461 555 BY SIMILARITY. 634 685 BY SIMILARITY. 638 BY SIMILARITY. 639 885 BY SIMILARITY. 631 1323 BY SIMILARITY. 631 1323 BY SIMILARITY. 632 1323 BY SIMILARITY. 634 1469 BY SIMILARITY. 635 55 POTENTIAR. 636 POTENTIAL. 637 891 POTENTIAL. 640 POTENTIAL. 651 600 POTENTIAL. 652 600 POTENTIAL. 653 993 POTENTIAL. 663 893 POTENTIAL. 664 694 POTENTIAL. 676 696 POTENTIAL. 676 696 POTENTIAL. 677 697 POTENTIAL. 677 697 POTENTIAL. 678 697 POTENTIAL. 678 697 POTENTIAL. 679 697 POTENTIAL. 670 697 POTENTIAL. | 5.5%; Score 661; DB 1; Length 1476; ilarity 24.1%; Pred. No. 3.83e-116; Conservative 407; Mismatches 612; Indels 140; Gaps 112 | <pre>dskymvlvpsqlytetpekiclhlyglnetvtvtaslvsqsgrknlfde-lv-ldkdl-f 84 : :: : :: : :: : : EQTYVISAPKIFRVGASENIVIQVYCYTEAFDATISIKSYPDKKFYSSGHVHLSSENKF 79</pre> | <pre>qcvsfi-i-pr-lsssdee-dflyvdikgpthefekrkavlvknkesvvfvqtdkpvykp 140 ::: : ::: : : : : : : : </pre> | gqsvkfrvvsmdkmlrplnellplayiedpkknri-mqwrdiktenglkqmsfslaaepi 199 : : : : : | <pre>ggpykivv-hkesgekeehs-ftvmefvlprfnvdlkvp-namsvnd-evlsvtacgkyt 255 </pre> | ygkpvp-ghvkinv-cret-etgcrev-nsqldnng-cs-tqevni-telqskk-rnyev 307 : : : : : : | q-lf-hvnatvteegtgl-eførsgttkieritnkliflkadshf-rhgipffvkvr 360 : | lvdikgdpipnekvfikagelsytsatttdqhglaefsidttcisgsslhikvnhkeeds 420 : :: :: :: : VKDSLDQLVGGVPVIINAQTIDVNQETSDLDPSKSVTRVDDG-VASFVLNLPSGVTVLEF 433 | csyfycmeerhasakhvay-av-ys-lsksyiyld-tetssilpcnqihtvqahfilkgd 476 : : : | <pre>lgvlkelifyylvmaggsligtgnhthqvepgeapvkgkfaleipvefsmvpmakmliyt 536 : : : : : : x-iDKITHYNYLILSKGKIHFGTRE-KFSDASYQSINIPVTQNMVPSSRLLVYY 544</pre> |
|--|--|--|--|---|---|--|--|--|---|--|
| 22 20 10 10 10 10 10 10 10 10 10 10 10 10 10 | Simil 68; | dskymvlvps : :: EQTYVISAPK | qcvsfi-i-F ::: QNSAILTIQP | gqsvkfrvve | qqpykivv-h YGMWTIKAKY | ygkpvp-ghv : YNKVVTEADV | q-1f-hv : | lvdikgdpip : :: VKDSLDQLVG | csyfycmeer : NVKTDAPDLP | lgvlkelify : : Y-IDKITHYN |
| FT DISULFID FT DISULFID FT DISULFID FT DISULFID FT DISULFID FT DISULFID FT DISULFID FT DISULFID FT CARBOHYD FT CARBOHYD | Query Match Best Local Matches 3 | 0b 28 Qy 20 | Db 85 Qy 80 | Db 141 Qy 138 | Db 200 Qy 196 | Db 256 Qy 256 | Db 308 Qy 316 | 0b 361 Qy 375 | 0b 421 Qy 434 | Db 477 Qy 492 |
| | | n o | <u>п</u> о | Δ Ο | <u>п</u> Ф | 0 0 | Δ Φ | 0 0 | <u> </u> | ы O |

112;

| Jan 28 12.23 | US:08-487-285.A.2.rsp | 47 |
|----------------|--|------------|
| · | | 590 |
| 545 I 591 a | 545 IVTGEQTAELVSDSVALNIEEXC-GNQLQVHLSPDADAYSPCGIVSLANATGADSWVALA 6 591 avdasvillknessisneswivnibomacmkfvnssriseddeddiusswiaekhtnivn 6 | 603 650 |
| − ₹ | | 663 |
| Ġ | 651 hgtekdvyryved-mg-lta-ftnlmiklpiicfdygmv-pisaprvefd 6 | 969 |
| 12 | 664 ENDEPCKEILRPRATLØKKIEELAAKYKHSVYKKCCYDGACVNNDETCEGRAARISLGPR 7 | 723 |
| 1 | 697 -l-aftpeiswslrttlskrpeepprkdpssndpltet-irkyfpetwywdivtynst 7 | 751 |
| O | , HLVPRR | 783 |
| 752 g | glaevemtvpdtitewkagalclsndtglglssvvplqafkpffvevslpysvvrgeafm 8 | 811 |
| 784 - | | 840 |
| 812 1 | | 698 |
| 841 L | | 895 |
| 870 v | | 927 |
| N 968 | SYSGVTLDPRGIYG | 951 |
| 928 e | | 978 |
| 952 T | | 1011 |
| 日 - | 979 mvlfapniyvlkyln-etq-qltqkiktka-lgf-lragygrelnykhkdgsysafgd 1 | 1032 |
| - 🗔 | TGNHWNIFHSDPLIEKQKLKKKLKEGMLSIMSYRNADYSYSVW | 1069 |
| ъ. | arafifideshithaftwlsqk-qkdngcfrssgslfnn | 1091 |
| | : :: : : :: :: : : : : | 1126 |
| 1092 a | a-mkggvd-demtleayit-mallessip-athpvvskalsclesswktiegernasfvy 1 | 1147 |
| Н | 1127 IKLQCTLPVEARENSLYLTAFTVIGIRKAFDICPLVKIDTALIKADNFLLENTLPAQSTF 1 | 1186 |
| - ب | | 1205 |
| 1187 T | TIAISAYALSLGDKTHPQFRSIVSALKREALVKGNPPIYRFWKDNLQHKDSSVPNTGT 1 | 1244 |
| м - | | 1264 |
| - Z | 1245 ARMVETTAYA-LLTSLNIKDINYVNPVIKMISEEQRYGGGFYSTQDTINAIEGLT 1 | 1298 |

1323 cvyaq-tmlrynmhlekqisafaiwvqtvpltcnnpkqh-ns-fqisle-isytgsr--p 1376 : | : : : | | | : | | | : | 136 | 1356 IATVHVTTVVHKTSTSEEVGSFYIKIDTQDIEASHYRGYGNSDYKRIVACASYKPSREES 1415

Jan 28 12:23

US-08-487-283A-2.rsp

\$

Db 1432 sfiiqqdipvrnlqpaivkvydyyetd 1458

Qy 1476 RFRIFELEEVGFLSPATFTVYEYHRPD 1502

Search completed: Wed Jan 28 12:17:15 1998 Job time : 69 secs.

US-08-487-283A-2.rag

Jun 28 12:20

| | (TM) |
|---------|---------|
| | |
| | _ |
| <u></u> | - |
| | : :- |
| | |
| | := |
| | |
| | |

Release 2.1D John F. Collins, Biocomputing Research Unit. Copyright (c) 1993, 1994, 1995 University of Edinburgh, U.K. Distribution rights by IntelliGenetics, Inc.

protein - protein database search, using Smith-Waterman algorithm MPsrch_pp

Wed Jan 28 12:12:44 1998; MasPar time 34.21 Seconds 602.393 Million cell updates/sec Run on:

Tabular output not generated.

>US-08-487-283A-2 (1-1676) from USO8487283A.pep 12048

Perfect Score: Description:

1 MGLLGILCFLIFLGKTWGQE..........CQAFLANLDEFAEDIFLNGC 1676 Sequence:

Scoring table:

PAM 150 Gap 11

101610 seqs, 12294212 residues Searched:

Post-processing: Minimum Match 0% Listing first 45 summaries

i.parti 2:part2 3:part3 4:part4 5:part5 6:part6 7:part7 8:part8 9:part9 10:part10 11:part11 12:part12 13:part13 14:part14 15:part15 16:part16 17:part17 18:part18 19:part19 20:part20 21:part21 a-geneseq28 Database:

Mean 40.813; Variance 192.520; scale 0.212

Statistics:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

| 12 | 540 | 4.5 | 403 1 | <u>س</u> ج | R71933 | Cobra venom factor C- | 6.15e-34 |
|----|-----|------|--------|------------|--------|-----------------------|------------|
| 14 | 436 | י נפ | 344 | r ~ | R10899 | Rat phospholipase A2 | 4.57e-25 |
| 15 | 429 | 3.6 | | , 0 | R51948 | pholipase A2 i | 1.78e-24 |
| 16 | 438 | 3.6 | | 2 | R51949 | | 3.10e-25 |
| 11 | 438 | 3.6 | 349 | 4 | R21776 | Phospholipase A2 inhi | 3.10e-25 |
| 18 | 438 | 3.6 | | m | R10900 | Human phospholipase A | 3.10e-25 |
| 19 | 141 | 1.2 | | 15 | R77605 | Pro-C5 polypeptide KS | 1.42e - 01 |
| 20 | 139 | 1.2 | | 7 | P71532 | Sequence of new pepti | 1.96e-01 |
| 21 | 116 | 1.0 | 23 | ~ | P71534 | Sequence of new pepti | 6.86e+00 |
| 22 | 105 | 0.9 | | 7 | P71533 | of new | 3.43e+01 |
| 23 | 105 | 6.0 | 153 1 | | R60520 | Human alpha-2-MRBDv. | 3.43e+01 |
| 24 | 109 | 6.0 | | بو | R94906 | RAP-1 radiation prote | 1.93e+01 |
| 52 | 110 | 6.0 | | 12 | R60882 | P. abyssi DNA polymer | 1.67e+01 |
| 56 | 104 | 0.9 | | 2 | R60883 | P. occultum DNA polym | 3.96e+01 |
| 23 | 104 | 6.0 | | ب | R92523 | Pyrodictium occultum | 3.96e+01 |
| 28 | 110 | 0.9 | 903 1 | ب | R87007 | Hyperthermostable pro | 1.67e+01 |
| 53 | 110 | 6.0 | | ٠ | R87008 | Protease. | 1.67e+01 |
| 30 | 100 | 8.0 | | 7 | R11547 | T. hyo 39 kD family 2 | 6.98e+01 |
| 31 | 102 | 8. | | =: | W20797 | H. pylori transporter | 5.27e+01 |
| 32 | 100 | 9.0 | | 11 | R88328 | Cotton UDP glucose sy | 6.98e+01 |
| 33 | 96 | 8.0 | 522 1 | 6 | W06330 | 7 | 1.22e+02 |
| 34 | 101 | 8.0 | | ₹. | R74187 | Chick p75. | 6.06e+01 |
| 32 | 101 | 0.8 | | 15 | R90696 | Borrelia burgdorferi | 6.06e+01 |
| 36 | 101 | 8.0 | | S. | R90699 | Borrelia burgdorferi | 6.06e+01 |
| 31 | 100 | 8.0 | 628 1 | ~ | R64979 | Pig Syk. | 6.98e+01 |
| 38 | 101 | 8.0 | | ₹. | R72737 | Plasmodium falciparum | 6.06e+01 |
| 39 | 96 | 8.0 | | م | W06331 | Full-length receptor- | 1.22e+02 |
| 40 | 96 | 0.8 | | 4 | R85930 | Protein tyrosine-kina | 1.22e+02 |
| 41 | 96 | 8.0 | | ن | W06335 | Full length receptor- | 1.22e+02 |
| 42 | 96 | 0.8 | | 0 | W11304 | Receptor-type tyrosin | 1.22e+02 |
| 43 | 86 | 8.0 | | S | R29648 | AmEPV Spheroidin prot | 9.22e+01 |
| 44 | 86 | 8.0 | | 0 | R55576 | AmEPV spheroidin. | 9.22e+01 |
| 45 | 86 | 8.0 | 2199 1 | ٠ | R94562 | Human cytotactin. | 9.22e+01 |
| | | | | | | | |

ALI GNMENTS

m

| Db 481 hlniivtękspyidkithynylilskgkiihfgtrekfsdasygsinipvtgmmypssrl 540 | 541 | Qy 541 LVYYIVTGEQTAELVSDSVWIANIEEKCGNQLQVHLSPDADAYSPQGTVSLAMATGMDSWV 600 | 601 alaavdsavygvqrgakkplervfqfleksdlqcqagglnnanvfhlagltfltnanad | 601 ALAAVDSAVYGVQRGAKKPLERVFQFLEKSDLGCGAGGLINNANVFHLAGLTFLTNANAD | 661 dsqendepckeilrprrtlqkkieeiaakykhsvvkkccydgacvnndetceqraarisl | DSQENDEPCKEI LRPRRTLQKKI EE I AAKYKHSVVKKCCYDGACVNNDETCEQRAAR I SL | | (| Db 841 lkgtvynyrtsgmqfcvkmsavegictsespvidhggtksskcvrqkvegssshlvtftv 900 | Db 901 lpleiglhninfsletwfgkeilvktlrvvpegvkresysgvtldprgiygtiørrkefp 960 | Db 961 yripldlypkteikrilsvkgllygeilsavlsqeginilthlpkgsaeaelmsvypvfy 1020 | Db 1021 vfhyletgnhwnifhsdpliekqklkkklkegmlsimsyrnadysysvwkggsastwlta 1080 | Db 1081 falrulgqvnkyveqnqnsicnsllwlvenyqldngsfkensqyqpiklggtlpvearen 1140 | Db 1141 slyltaftvigirkafdicplvkidtalikadnfllentlpaqstftlaisayalslgdk 1200 | Db 1201 thpqfrsivsalkrealvkgnppiyrfwkdnlqhkdssvpntgtarmvettayallteln 1260 | <pre>Db 1261 lkdinyvnpvikwlseeqrygggfystqdtinaieglteysllvkqlrlsmdidvsykhk 1320 </pre> | Db 1321 galhnykmtdknflgrpvevllnddiivstgfgsglatyhvttvvhktstseevcsfylk 1380 | Db 1381 idtqdieashyrgygnsdykrivacasykpsreesssgsshavmdislptgisaneedlk 1440 |
|---|-----|--|---|--|--|--|--|------------------------|---|---|--|---|---|---|---|---|---|---|
| site | | /label= N-glycosylation_site Oy | 005688. US-236208. | er EE, Nye SH, Rollins S; | Thomas TC; | plement C5 | Ine CLNA sequence of the compliament to gene transcript predicts a secreted pro-C5 precursor of 1576 amino acids (R71504). C5 is a beta-c1obulin heterodimer thought to play a rate in the pathogenesis Db | na-chain Monoclonal | and humanised recombinant antibodies that recognise the alpha-chain KSSKC epitope (R7765) block C5a generation, thereby reducing glomerular inflammation and kidney dysfunction associated with GN. Sequence 1676 AA; | Query Match 100.0%; Score 12048; DB 15; Length 1676; Best Local Similarity 100.0%; Pred. No. 0.00e+00; Matches 1676; Conservative 0; Mismatches 0; Indels 0; Gaps 0; Qy | <pre>1 mgllgilcfliflgktwgqeqtyvisapkifrvgaseniviqvygyteafdatisiksyp 60 1 </pre> | 61 dktfsyssghvhlssenkfgnsailtigpkqlpggppvsyvlevvskhfskskrmpit 120 | 121 ydngflfihtdkpvytpdgsvkvrvyslnddlkpakretvlfidpegsevdmveeidhi 180 Db | 181 giisfpdfkipsnprygmwtikakykedfsttgtayfevkeyvlphfsvsiepeynflgy 240 Db | 241 knfknfeitikaryfynkvyteadvyitfgiredlkddqkemmqtamqntmlingiaqvt 300 | 301 fdsetavkeløyysledinnkylyjavtviestggfseeaeipgikyvløpyklnlvatp 360 | 361 lflkpgipypikvqvkdsldqlvggvpvilnaqtidvnqetsdldpsksvtrvddgvasf 420 | 421 vinipsgytviefnyktdapdlpeenqaregyraiaysslagsylyidwtdnhkallvge 480 |

В

g

õ

셤

ð g ð Б õ

ð

g

Š

셤 8

ð

Š

Ø

셤 ð 음 g g 셤 g a 셤 용 à ð ð Š ð 8 ò 1441 alvagvdqlftdyqikdqhvilqlnsipssdflcvrfrifelfevqflspatftvyeyhr 1500 1441 ALVECVDQLFTDYQIKDGHVILQLASIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHR 1500 pdkqctmfystsnikiqkvcegaackcveadcgqmqeeldltisaetrkqtackpeiaya 1560 ykvsitsitvenvfvkykatlldiyktgeavaekdseitfikkvtctnaelvkgrqylim 1620 X. laevis; pre-pro molecule; beta chain; alpha chain; codon usage; Pre-pro-cobra C3 protein. Cobra; C3; third component of complement; human; mouse; rat; G+C content; immune response; host defence; glycosylation; tertiary structure; maturation; thioester site; Fritzinger DC, Vogel C; /note= "Potential glycosylation site" /note= "Potential glycosylation site" /note= "Potential glycosylation site" Protein 739..1269 'note≂ "Potential glycosylation site" Location/Qualifiers Binding_site 1214..1219 /note= "Potential CR2 binding site" LT 2 R63222 standard; Protein; 1651 AA. Binding_site 1419..1442 /note= "Properdin binding site" alternative pathway activation Binding_site 1176..1201 /note= "Factor H binding site" Binding site 1221..1235 /note= "Factor H binding site" Binding_site 742..751 /note= "Factor B binding site" 1176..1201 30-JUN-1995 (first entry) 1270..1651 996..1017 1..22 "Signal sequence" Active_site 996..101 /note="Thioester site" 07-APR-1993; US-043747. (GEOU) UNIV GEORGETOWN. 23..655 'note= "Alpha-chain" 1352 158 Protein /note= "Gamma-chain" 07-APR-1994; U03441. Protein 12 /note= "Beta-chain Modified site Modified_site Modified site Modified site Bredehorst R, /note= "Prop W09423024-A. 3-0CT-1994. Naja naja. Protein Peptide /note= 1501 1561 1621 1621 1561 요 ð 유 a 음 ð ð ð

CVF 1 and CVF 2 - which are used in the DNA encoding cobra C3, treatment of cancer Jan 28 12:20

Claim 1; Fig 2A-2L; 155pp; English.

The cDNA sequence of cobra C3 shows a high sequence homology with C3 This sequence represents cobra C3 (third component of complement)

molecules from human, mouse, rat and X. Jaevie. Cobra C3 is synthesised as a pre-pro molecule that is subsequently processed into the mature two-chain protein by removing the signal peptide and the four Arg residues between the beta and alpha chain. The alpha chain comprises 992 amino acids and the beta chain comprises 633

similarity in the tertiary structure of mammalian and cobra C3. There are four Arg residues in pre-pro-C3 which are required for the proper maturation to yeild the secreted C3 molecule, and the thioester site, contrast to C3 molecules studied so far, cobra C3 is not glycosylated. All 27 Cys residues are conserved indicating a very high degree of residues, being 12 residues shorter than the human beta chain. In

which is unique to the alpha chain, is required for alternative pathway different codon usage compared to mammalian C3 mRNAs. The G+C content of all known mammalian C3 mRNAs is more than 53%. The G+C content of cobra C3 mRNA is significantly lower at 43%. The significance of this difference is not known. C3 is thought to have important functions in activation and covalent binding to target cells. Cobra C3 has a the immune response and host defence.

Sequence

98; Gaps 518; Conservative 416; Mismatches 655; Indels 127; Length 1651; 19.7%; Score 2369; DB 12; 30.2%; Pred. No. 1.94e-197; Best Local Similarity Query Match Matches

9 malylvaalligfpg-s-shgalytlitpavlrtdteeqilveahgdstpksldifvhdf : :

1 MGLLGILCFLI-FLGKTWGQEQTYVISAPKIFRVGASENIVIQVYGYTEAFDATISIKSY

prkqktlfqsrvdmnqagsmfvtpti-kvpakelnkdskqnqyvvvkvtgpqvalekvvl 120 62

60 PDKKFSYSSGHVHLS-SENKFQNSAILTIQPKQI,PGGQNPVSYVYLEVVSKHFSKSKRMP 118 --..

psgsir-p-ynlpelvsfgtwkavakyehspeesytayfdvreyvlpsfevrlgpsdkfl 237 ----

-yidgnknfhvsitarylygkkv-egvafvvfgvk--i-ddakksipdsltriplidgdg 292 238

239 GYKN-FKNFEITIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLINGIA 297

293 eatlkrdt-1rs-rfqdlnqlvghtlyvsvtvitesgsdmvvteqggihivtspyqiyft 350 351 ktpkyfkpgmpyeltvyvtnp-d---gs-paa-hvp-v-vs-ea--ih-segtt-lødgt 397

398 aklilntplniqslpitvrtnhgdlprerqaiksmtatayqtqggsenylhvaitsteik 457

418 ASFVLALPSGVTVLEFNVKTDAPDLPEENQAREGYRAIAYSSLSQSYLYIDWTDNHKALL 477

pgdnlpvnfnvrgnanslnqikyftylilnkgkifkvgrqprrdgqnlvtmnlhitpdli 517 458

요

ð

g

8 & B

g

õ

셤

Š

g

õ

g

Š

g

õ

g

à

셤

S S

Š

g

õ

a

8

g

õ

Š

~

ctry-1g--evds-tm-tiidismltgffpdaedlkrlsngvdryiskfeidnnmaqkgt 1431

1377

셤

Š

Jan 28 12:20 US-08-487:283A-2.rag

00

| 6y | : : : ::: : : ::: : | 55 |
|----------------|---|----|
| QQ | 1432 vviyldkvshsedeclhfkihkhfevgfiqpgsvkvysyynldeqckfyhpdketgvln 1491 | 5 |
| δ | 1460 VILQINSIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHRPDKQCTMFYSTSNIK-I-Q 1517 | Ξ |
| 임 | kichgnicrcaeetcsllnqq-k-kidlqlriqkacaqnvdyvyktkllrieekdgndiy : : : | 2, |
| δλ | EELDLT I SAETRKQTACKP EI AYA | Ε. |
| qq | 1550 fmdvlevikggtdrnaqakarqyvsqrkcqealnlkldndyliwglssdlwpmk-d-dis 1607 | 6 |
| δy | KIGEAVAEKDSEITFIKKVTCTNA-ELVKGRQYLIMG-KEAL-Q | 34 |
| В | 1608 ylit-kn-twierwpnedecqd-eef-qnlcddfaq 1639 | |
| ớ | 1635 YIYPLDSLTWIEYWPRDTTCSSCQAFLANL-DEFAE 1669 | |
| RESULT ID R | III 3 BAGNAN etandard: Drotein: 1663 MA | |
| AC : | | |
| Į i | 2 | |
| 1 Z | numan mournied of (1735), 1735, 1736). C3 protein; convertase; Factor I; Factor H; complement; tumour; | |
| 3 8 | | |
| 3 % | Symplectic. Key Location/Oualifiers | |
| 딥 | tide | |
| E 6 | /label= Sig_peptide Protein 73667 | |
| FT | "C3 beta | |
| E 8 | 668671 | |
| - t- | <u>,</u> & | |
| : E | ains" | |
| FI | Protein 6721663 | |
| FT | 급 | |
| N G | MO9607/38-AZ | |
| 7. F | 14-mars - 1996. 08-SEP-1995: G02121. | |
| PR | 08-SEP-1994; GB-018147. | |
| PR | 04-MAY-1995; GB-009102. | |
| PA DI | (IMUT-) IMUTRAN LTD. | |
| DR | Farines 10, national Ma; WPI; 96-171613/17. | |
| PT | protein forming stable C | |
| Ld d | for generalised complement depletion or localised complement | |
| PS | dctrvalion Claim 11; Fig 1; 81pp; English. | |
| ខ | A modified human C3 protein (R94030) differs from the wild-type | |
| ខ | (R94028) by substitution of Asp-Glu-Asp at positions 752-754 by | |
| 3 8 | GLY-Ser-GLY. It is obtained by site-directed mutagenesis of C3-encoding cDNA (117738). The modification reduces the | |
| ខូខ | interaction of C3b/C3i with Factor H in comparison to wild-type | |
| ပ္ပ | C3. This allows the modified C3 to be used therapeutically to | |
| 38 | super-active the complement system or the increase a target's (e.g. tumour, bathogen or virus-infected cell) sensitivity to | |
| ខ | nt-mediated destruction. | |
| ος | Sequence 1663 AA; | |
| Ö | Query Match 19.4%; Score 2339; DB 16; Length 1663; | |
| í | THITTHEFTE SOLOW, FIGURE NO. 1 | |

| 9 | ۰ | 8 | ÷ | ŀ, | R |
|---|----|---|---|----|----|
| ٠ | | ۰ | ٠ | ۰ | ٥ |
| : | × | х | ٧ | ٠ | В |
| ٠ | ۰ | ۰ | ۰ | ٥ | ٥ |
| ۶ | þ | з | × | ۹ | × |
| ٠ | ٠ | ۰ | ۰ | ۶ | ٥ |
| ۶ | | 3 | ŏ | ı | 8 |
| ٠ | ۰ | ۰ | ۰ | ۶ | ¢ |
| ٠ | ٠. | 8 | и | × | s |
| è | ٠ | ۰ | ۰ | ۶ | ٥ |
| 9 | | s | ٥ | 3 | ٥ |
| 2 | 3 | ٠ | ٠ | ı | e. |
| ٠ | ٠ | ۰ | ۰ | 8 | ٥ |
| ۶ | | з | × | ٠ | 8 |
| ٠ | ۰ | ۰ | ۰ | b | ٥ |
| 5 | 9 | я | ٠ | ŀ. | 8 |
| ٠ | ٠ | o | ۰ | 2 | ø |
| ٠ | | × | ч | r. | ٥ |
| ۰ | и | ٥ | ٥ | × | × |
| ٠ | ٠ | ۰ | ۰ | ٠ | ٥ |
| > | | ٥ | × | × | × |
| ٠ | ۰ | ۰ | ۰ | ٥ | 9 |
| : | ŀ | 8 | х | ٠ | В |
| ٠ | ٠ | 8 | ٥ | ١ | e |
| | × | × | × | 8 | 8 |
| ď | × | 8 | × | 9 | 8 |
| ٠ | | 8 | × | 8 | 8 |
| è | × | 8 | × | S | × |
| ٤ | ٥ | ۲ | ٥ | ٠ | ì |
| ۶ | i | × | ď | 'n | S |
| ٠ | ٠ | ٧ | 1 | 7 | e |
| ۶ | ı | ø | ı | r. | 8 |
| ٥ | · | è | ú | S | ó |
| ۶ | ۶ | ١ | ĕ | 2 | 8 |
| è | ۰ | ď | 3 | í | é |
| 4 | ĸ | ٠ | ŕ | í | ĕ |
| ٠ | 3 | ķ | ٠ | Š | S |
| ¢ | ٤ | è | ٠ | ŀ | í |
| 3 | 3 | ч | | ì, | 8 |
| ò | | 1 | 4 | ĸ | 8 |
| ı | F | 9 | z | Ŋ | ٥ |
| 1 | ٠, | r | ٩ | r. | К |
| c | • | 3 | c | 2 | ٥ |
| 1 | я | | н | ĸ | К |
| ٠ | 7 | ٠ | 6 | r. | Ŋ. |
| 2 | £ | | | ٥ | ٥ |
| 2 | | | | н | |
| | | | | | |
| 1 | 1 | ė | ٠ | ì | ٥ |
| į | Ì | Ċ | ۶ | ì | ì |
| | Ì | į | į | ١ | |
| | ١ | į | į | | |
| | • | į | Ì | | |
| | | į | Ì | | |
| | | į | | | |
| | | | | | |
| | | | 3 | | |
| | | | | | |
| | | | 3 | | |
| • | | | 2 | | |
| • | | | 3 | | |
| | | | 3 | | |
| | | | 3 | | |
| | | | 3 | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| * | | | 3 | | |
| * | | | | | |
| * | | | | | |
| ** | | | | | |
| *** | | | | | |
| *** | | | | | |
| (, | | | | | |
| ((() | | | | | |
| (() | | | | | |
| *** | | | | | |
| *** | | | | | |
| ** | | | | | |
| ** | | | 3 | | |
| **** | | | | | |
| ** | | | | | |
| *** | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |

Jan

o,

| Mat | Matches 488; Conservative 459; Mismatches 660; Indels 115; Gaps 90; | δ |
|--------|--|----------|
| a | œ | q |
| ò | : : : : : | Qy |
| 원 상 | 65 kklvissektvitpatnhmgnvtftipanrefksekgrnkfvtvqatfgtqvve-kvvlv 123 | Db Qy |
| අු | o 124 sląsgylfiqtdktiytpgstylyriftvnhkllpvgrtvmvnienpegipykqdslssq 183 | qu |
| ģ | 120 | δ |
| ପୁ |) 184 nqlqvlpls-wdipelvnmqqwkirayyenspqqvfstefevkeyvlpsfeviveptekf 242 | qu |
| ò | 178 DHIGIISFPDFKIPSNPRYGMWTIKAKYKEDFSTTG | Qy |
| QQ | 243 yyi-yne-kglevtitarflygkkv-egtafvifgi-qdgegrislpeslkripiedg 296 | qu |
| ð | | λö |
| ପ୍ପ | 297 sgevvlørkvlldgvqnpraedlvgkslyvsatvilhsgsdmvqaersgipivtspyqih 356 | qa |
| ò | 296 IAQVTEDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSE | δλ |
| QQ | | qu |
| ŝ | 356 LVATPLEIKPGIPYPIKVQVKDSIDQIVGCVPVIINAQTIDVNQETSDIDPSKSVTRVDD 415 | Qy |
| qq | • | qu |
| δ | 416 GVASFVINLPSGVTVILEFNVKTDAPDILPEENQAREGY | Qy |
| d d | | QO |
| 6ý | 476 LLVG | Qy |
| QQ | | qq |
| ó | 534 MVPSSRLLVYYIVTGEQTA-ELVSDSVWINIEEKCGNQLQVHLSPDADAYS-PGQTVSIN 591 | Qy |
| g Q | 585 iegdhgarvvlvavdkgvfvlnkknkltgskiwdvvekadigctpgsgkdyagvfedagl 644 | qa |
| δy | 592 | θy |
| g | 645 | qu |
| ολ | 652 | . Qy |
| g | 703 mrfscgrrtrfislgeackkvfldccnyitelrrqharashlglarsnlgsgiiaeeniv 762 | QQ |
| ٥y | 708 DE-TCEQRAARISLGPRCIKAFTECCVVASQLRANISH-KD | ρ |
| qq | 763 grsefpeswlwnvedlkeppkngistklmniflkdsittweilavsmsdkgicvadpfe 822 | RE |
| δ | 765 IRSYFPESMIMEVH-L-VPRRKQI-QFALPDSLTTWEIQGIGISNT | 10 |
| q | 823 | DT |
| δλ | 817 AKVFKD | KA. |
| ପ୍ର | 883 qqtitippksslavpyvivplktglqevevkaavyhhfisdqvrkslkvvpeg 935 | FH |
| | | |

Jan 28 12:20 US-08-487-283A-2 rag 10

| Ś | ZY 0/0 VOINSSNOVNKNYEGSSSNEVIFIVEFELGENNINFSEEIMFGNEIL | UI TUANEER | 222 |
|----------|--|--------------------|------|
| qu | 936 | gtpvagmte | 366 |
| 0y | 934 | CLLVGEILS | 686 |
| QO | 993 davdaerlkhlivtpsgcgegnmigmtptviavhyldete | krqgaleli | 1049 |
| δy | 066 | EKQKLKKKL | 1049 |
| q | Db 1050 kkgytqqlafrqpssafaafvkrapstwltayvvkvfslavnliaidsqvlcgavkwlil | cgavkwlil | 1109 |
| δλ | 1050 | CNSLLWLVE | 1109 |
| DP | 1110 | eqvnslpgs | 1168 |
| δy | 1110 | -LVK-IDTA | 1167 |
| qu | Db 1169 itkagdfleanymnlqrsytvaiagyalaqmgrlkgpllnk-flttakdk-nr- | takdk-nr- | 1219 |
| ργ | 1168 | KGNPPIYRF | 1227 |
| qa | 1220 | yygggygst | 1269 |
| δy | Δ) 1228 WKDNLQHKDSSVPNTCTARAVETTAYALLTSIALKDINYVRPVIKWISEEΩRYGGGFYST | RYGGGFYST | 1287 |
| qa | 1270 | seet keneg | 1329 |
| Qy | 2) 1288 QDTINAIEGLTEYSL-LVKQLRLSMDIDVSYKHKGALHNYKMT-DKNFLGRPVEVLINDD | PVEVLLNDD | 1345 |
| ΩP | 1330 | lakntmilei | 1388 |
| δy | 2) 1346 LIVSTGFGSGLATVHVTTVHKTSTSEEVCS-FYLKIDTQDIEASHYRGYGNSDYKRIVA | : : : NSDYKRIVA | 1404 |
| 셤 | 1389 | dkafsdrnt | 1443 |
| Qy | : : :: :: :: : : : : :: : : | -KD-GH- | 1459 |
| qq | 1444 | pekedgkln | 1503 |
| Qy | 2y 1460 VILQIANSIPSSDFLCVRFRIFELEEVGFLSPATFTVYEYHRPDKQCTMFY-S-TSNIKIQ | S-TSNIKIQ | 1517 |
| OP | 1504 | qlsndfdey | 1561 |
| δy | 1518 | TVENVEVKY | 1577 |
| qa | Ob 1562 imaieqtiksgsdevqvgqqrtfispikcrealkleekkhylmwglss-df-wgekpnls | -wgekpnls | 1619 |
| . 03 | 1578 | KYNFSFRYI | 1636 |
| qq | | | |
| Qy | 2y 1637 YPLDSLTWIEYWPRDTTCSSCQAF-L-ANLDEFAEDIFINGC 1676 | | |
| REST | RESULT 4 | | |
| 0 % | | | |
| TO | 21-MAY-19 | | |
| X. | | | |
| OS FH | Homo sapiens. | | |
| FT | | | |

Jan 28 12:20

405 525 645 763 285 셤 8 g ð В 8 g ð 셤 ð 유 ò g ð g ò g 8 g 유 g ð g ð 쇰 ð 염 8 음 õ ð 9 297 sgevvlsrkvlldgvqnpraedlvgkslyvsatvilhsgsdmvqaersgipivrspyqih 356 ::| : ||| ||| ||: ||| :|| :|| || ||| :296 IAQVTFDSETAVKELSYYSLEDLNNKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLN 355 62 KKESYSSGHVHLSSENKFQNSAILTIQP-KQLPGGQNPVSYVYLEVV-SKHFSKSKRMPI 119 124 slqsgylfiqtdktiytpgstvlyriftvnhkllpvgrtvmvnienpegipvkqdslssq 183 :::::|{|:||| :|| :| |::::| | | ::::| | | :: 243 yyi-yne-kglevtitarflygkkv-egtafvifgi-qd--geqrislpeslkripiedg 296 kklvlssektvltpatnhmgnvtftipanrefksekgrnkfvtvqatfgtqvve-kvvlv 123 nqlgvlpls-wdipelvnmgqwkirayyenspqqvfstefevkeyvlpsfeviveptekf 242 236 NFIGYKNFKNFEITIKARYFYNKVVTEADVYITFGIREDLKDDQKEMMQTAMQNTMLING 295 ftktpkyfkpgmpfdlmvfvtnp-d---gs-pay-rvp-vav-q--ge-dtvqsltq-gd 404 8 sll-ll-llthlplalg-spmysiitpnilrleseetmvleahdaggdvpvtvtvhdfpg 64 Gaps is a complement pathway protein that is suceptible to cleavage ant complement pathway protein forming stable C3 convertase generalised complement depletion or localised complement Human C3 protein (R94028) was produced by expression of a cDNA 458; Mismatches 661; Indels 115; C3 is a complement pathway protein that is suceptible to clea by Factor I and is also susceptible to the inhibitory action Length 1663; produced by site-directed mutagenesis. These mutants can be localised super-activation at a specific target to increase the target's sensitivity to complement-mediated destruction. sequence (T17738) isolated from a human liver cDNA library. of Factor H. Mutants of C3 (R94029 and R94030) have been used to super-active the complement system, or to induce Match 19.4%; Score 2339; DB 16; Local Similarity 28.3%; Pred. No. 1.02e-194; Peptide 668..671 /note= "amino acids 668-671 are removed when precursor is cleaved into the alpha and Disclosure; Fig 1; 81pp; English. = = = = 488; Conservative Mutant complement pathway 672..1663 (IMUT-) IMUTRAN LTD. Farries TC, Harrison RA; WPI; 96-171613/17. 14-MAR-1996. 08-SEP-1995; G02121. 08-SEP-1994; GB-018147. 04-MAY-1995; GB-009102. 23..667 /note= "C3 alpha chain" 'note= "C3 beta chain' Sig_peptide Ξ N-PSDB; T17738 409607738-A2. beta chains" activation /label= ? Protein Sequence Query Match Protein 65 Matches 184 357

8 6 6 6 B

8 8 8 8 8 8

Š

1050 kkgytqqlafrqpssafaafvkrapstwltayvvkvfslavnllaidsqvlcgavkwlil 1109 1287 1270 qatfmvfqalaqyqkdapdhqelnldvslqlpsrsskithrihwesasllrseetkeneg 1329 993 davdaerlkhlivtpsgcgeqmmigmtptviavhyldeteqwekf-g--lekrqgaleli 1049 1110 ekqkpdgvfqedapvihqemigglr-nnnekdmaltafvlislqeakdiceeqvnslpgs 1168 1169 itkagdfləanymnlqrsytvaiagyala--qmgr--lkg--pllnk-flttakdk-nr- 1219 1288 QDTINAIEGLTEYSL-LVKQLRLSMDIDVSYKHKGALHNYKMT-DKNFLGRPVEVLLNDD 1345 817 AKVFKDVFLEMNIPYSVVRGEQIQLKGTVYNYRTS-GMQFCVKMSAVEGICTSESPVIDH 875 935 gvaklsinthpsqkplsitvrtkkqelseaeqatrtmqalpystvgnsnnylhlsvlrte 464 416 GVASFVINIPSGVTVIEFNVKTDAPDIPEENQAREGYRAIAYSSISQSYLYIDWTDNHKA 475 465 lrpgetlnvnfllrmdraheakiryytylimnkgrllkagrqvrepgqdlvvlplsittd 524 fipsfrlvayytligasggrevvadsvwvdvkdscvgslvvksggsedrgpvpgggmtlk 584 644 651 tftsssggqtaqraelqcpqpaarrrrsvqltekrmdkvg-kypkel-rkccedgmrenp 702 703 mrfscgrrtrfislgeackkvfldccnyitelrrqharashlglarsnldediiaeeniv 762 srsefpeswlwnvedlkeppkngistklmniflkdsittweilavsmsdkkgicvadpfe 822 882 823 vtvmqdffidlrlpssvvrneqveiravlynyrqnqelkvrvellhnpafcslattkrrh | | |::: :| |||| ||:::: :|||| : : | 883 qqtit---ippk--ssls--vpyvivplktglqevevkaavyhhfisdgvrkslkvvpeg 1220 w-ed-pgkq--1-yn----veatsyallallqlkdfdfvppvvrwlnegryygggygst 1228 WKDNLQHKDSSVPNTGTARMVETTAYALLTSLNLKDINYVNPVIKWLSEEQRYGGGFYST ð

| g | | 1388 |
|----------|---|------|
| δ | 1346 LIVSTGFGSGLATVHVTTVVHKTSTSEEVCS-FYLKIDTQDIEASHYRGYGNSDYKRIVA | 1404 |
| Вр | 1389 ctryr-g-dgdat-msildismmtgfapdtddlkqlangvdryiskyeldkafsdrnt | 1443 |
| ò | 1405 CASYRPSREESSSGSSHAVMDISIPTGISANEEDIKALVEGVDQLFTDYQI-KD-GH- | 1459 |
| qq | 1444 liiyldkvshseddclafkvhqyfnveliqpgavkvysyynleestrfyhpekedgkln | 1503 |
| ŏ | 1460 VILQINSIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHRPDKQCTMFY-S-TSNIKIQ | 1517 |
| g | 1504 klcrdelcrcaeencf-iqksdd-kvtleerldkacepgvdyvyktrlvkvqlsndfdey | 1561 |
| ò | | 1577 |
| g | | 1619 |
| ò | 1578 KATLLDIYKTGEAVAEKDSEITFIKKVTCTNA-ELVKGRQYLIMGKEALQIKYNFSFRYI | 1636 |
| gg | 1620 yiigkdtwyehwpeedecqdeenqkqcqdlgaftesmyvfgc 1661 : : : : : : | |
| ò | 1637 YPLDSLTWIEYWPRDTTCSSCQAF-L-ANLDEFAEDIFLNGC 1676 | |
| RESULT | ULT 5 BOANOS etandard, Drotesin, 1663 Ah | |
| S S | R94029; | |
| H | 21-MAY-1996 (first entry) | |
| 음 3 | Ruman modified C3 (R1303X). C3 protein: convertace: Factor I: Factor H: complement: | |
| Ž | 11 | |
| ខ្លួ | Synthetic. | |
| E E | tide | |
| E | /label= Sig_peptide | |
| 5 | Protein 23667 | |
| : 5 | | |
| 뜶 | acids 668-671 are remo | |
| <u> </u> | precursor is cleaved into the alpha and beta chains" | |
| Ë | Protein 6721663 | |
| 딥 | /note= "C3 alpha chain" | |
| 5 | Misc difference 1303 | |
| Z Z | | |
| 2 | 14-MAR-1996. | |
| ᅜ | 08-SEP-1995; GUZ1Z1. 08-SEP-1994; GB-018147. | |
| 8 | 04-MAY-1995; GB-009102. | |
| PA D1 | (IMUT-) IMUTRAN LTD. | |
| z z | 17161 | |
| PT | t complement pathway protein formin | |
| 7 4 | for generalised complement depletion or localised complement | |
| S | ig 1; 81pp; English. | |
| ဗ ဗ | fied human C3 protein (R94029) differs from the very him substitution of Arc 1303 hu clutomic acid | |
| ខ | (nytoxo) by substitution of Arg-1905 by grundants actu, grytaine or glutamine. It is obtained by site-directed mutagenesis of | |
| ខ្ល | ONA (T17 | |
| 3 | resistance to cleavade by ractor 1 in comparison to wild-type to. | |

US-08-487-283A-2.rag Jan 28 12:20

90; 123 235 296 297 sgevvlsrkvlldgvqnpraedlvgkslyvsatvilhsgdmvqaersgipivtspyqih 356 ::| : : | | | | | | | | :: 295 296 IAQVTFDSETAVKELSYYSLEDIANKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKIM 355 357 ftktpkyfkpgmpfdlmvfvtnp-d---gs-pay-rvp-vav-q--ge-dtvqsltq-gd 404 gvaklsinthpsqkplsitvrtkkqelseaeqatrtmqalpystvgnsnnylhlsvlrte 464
||| : :| :: | :| :| :| :| :| :| :| 524 533 644 651 tftsssgqqtaqraelqcpqpaarrrrsvqltekrmdkvg-kypkel-rkccedgmrenp 702 707 mrfscgrrtrfislgeackkvfldccnyitelrrqharashlglarsnldediiaeeniv 762 64 108 DE-TCEQRAARISLGPRCIKAFTECCVVASQLRANISH-KDMQLGRLHMK-TLLPVSKPE 764 srsefpeswlwnvedlkeppkngistklmniflkdsittweilavsmsdkkgicvadpfe 822 Gaps 61 sll-ll-llthlplalg-spmysiitpnilrleseetmvleahdaggdvpvtvtvhdfpg kklvlssektvltpatnhmgnvtftipanrefksekgrnkfvtvqatfgtqvve-kvvlv nqlgvlpls-wdipelvnmgqwkirayyenspqqvfstefevkeyvlpsfeviveptekf 243 yyi-yne-kglevtitarflygkkv-egtafvifgi-qd--gegrislpeslkripiedg 316 NFIGYKNFRAFETIKARYFYNKVVTEADVYITFGIREDIKDDQKEMMQTAMQNTMLING lrpgetlnvnf1lrmdraheakiryytylimnkgr1lkagrqvrepgqdlvv1plsittd iegdhgarvvlvavdkgvfvlnkknkltgskiwdvvekadigctpgsgkdyagvfsdagl This allows the modified C3 to be used therapeutically to super-active the complement system or the increase a target's (e.g. tumour, pathogen or virus-infected cell) sensitivity to complement-mediated destruction. Score 2335, DB 16; Length 1663; Pred. No. 2.35e-194; 457; Mismatches 662; Indels 115; Ξ = --/ Match 19.4%; Local Similarity 28.3%; : - - :: = = 488; Conservative 1663 AA; Sequence Query Match 62 178 105 176 585 645 652 Matches 184 465 703 93 Best 88888 용 ð g 염 ð g ð 9 õ g рp g 셤 셤 В g 원 Š Š Š ò Š ð Š 9 à à

 g

õ

9

õ

ΩP

Š

g

Š

Jan 28 12:20

US-08-487-283A-2.rag

| *** | | | | | | | | | | | | | | | | |
|--|--|--|---|---|--|--|---|--|--|--|---|--|---|---|--|--|
| 2 | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | |
| | 816 | 882 875 | 935 933 | 992 | 1049 1049 | 1109 | 1168 1167 | 1219 1227 | 1269 1287 | 1329 1345 | 1388 1404 | 1443 1459 | 1503 1517 | 1561 1577 | 1619 1636 | |
| ###################################### | IRSYFPESWLMEVH-L-VPRRKQL-QFALPDSLTTWE1QGIGISNT-GICVADTVK | 823 vtvmqdffidLrlpssvvrneqveiravlynyrqnqelkvrvellhnpafcslattkrrh :::: ::::: ::: 817 AKVFKDVFIEMNIPYSVVRGEQIQLKGTVYNYRTS-GMQFCVKMSAVEGICTSESPVIDH | <pre>qqtitippksslsvpyvivplktglqevevkaavyhhfisdgvrkslkvvpeg : : : : :: ::: @QTKSSKCVRQKVEGSSSHLVTFTVLPLEIGLHNINFSLETWFGKEILVKTLRVVPEG</pre> | 936 irmnktvavrtldperl-gregvqkedip-padlsdqypdtesetril-lggtpvagmte :: : : | davdaerlkhlivtpsgcgegnmigmtptviavhyldeteqwekf-glekrggaleli | 1050 kkgytqqlafrqpssafaafvkrapstwltayvvkvfslavnliaidsqvlcgavkwlil ::: : : : : :: :: : : :: : 1050 KECMLSIMSYRNADYSYSVWKGCSASTWLTAFALRVLGQVNKYVEQNQNSICNSLLMLVE | 1110 ekqkpdgvfqedapvihqemigglr-nnnekdmalta_vlislqeakdiceeqvnslpgs | itkagdfleanymnlqrsytvaiagyalaqmgrlkgpllnk-flttakdk-nr- | w-ed-pgkql-ynveatsyallallqlkdfdfvppvvrwlnegryyggygst :: | <pre>qatfmvfqalaqyqkdapdhqelnldvslqlpsxsskithrihwesasllrseetkeneg : :: :: </pre> | <pre>ftvtae-gkgqqtlsvvtmyhakakdqltcnkfdlkvtikpapetekrpqdakntmilei : </pre> | <pre>ctryr-g-dqdat-msildismmtgfapdtddlkqlangvdryiskyeldkafsdrnt !: !: :: :</pre> | liiyldkvshseddclafkvhqyfnveliqpgavkvyayynleesctrfyhpekedgkln : : : :: : : : : : : : : | klcrdelcrcaeencf-iqksdd-kvtleerldkacerjydyyktrlvkvqlsndfdey : : : : | <pre>imaieqtiksgsdevqvgqqttfispikcrealkleekkhylmwglss-df-wgekpnls ::</pre> | yiigkdtwvehwpeedecqdeenqkqcqdlgaftesmvvfgc 1661 : : : : : : : : YPLDSLTWIEYWPRDTTCSSCQAF-L-ANLDEFAEDIFLNGC 1676 |
| 77.71 97 | 765 | 823 | 883 | 936 934 | 993 | 1050 1050 | 1110 | 1169 | 1220 | 1270 1288 | 1330 1346 | 1389 1405 | 1444 1460 | 1504 1518 | 1562 1578 | 1620 1637 |

RESULT

õ

ዋ

අ

දු දු

õ

Query Match 18.4%; Score 2213; DB 12; Length 1642; Best Local Similarity 29.3%; Pred. No. 2.66e-183; Matches 502; Conservative 438; Mismatches 642; Indels 132; Gaps 103; This sequence represents the cobra venom factor 1 (CVF1). The CVF1 protein is contained within cobra venom and is a structural analog of cobra G3 (third component of complement). It is thought that CVF1 and C3 are derived from different but realated genes. The thioseter site and factor H binding site of G3 and CVF1 are very similar, even though neither are actually present in the mature CVF1 protein. Factors H and an antibody which recognises a specific tumour cell. As CVF1 and CVF2 are insensitive to factor H control this method will lead to the I are able to regulate the activity of complement by dissociating C3b, Bb, and by cleaving C3b, CVFI is resistant to this regulation. CVFI 60 PDKKFSYSSGHVHLSSENKFQNSAILTIQPKQLPGGQNPVSYVYLEVVSKHFSKSKRMPI 119 122 sygssflíigtdkgiytpgspvlyrvísmdhntskmnktvivefgtpegilvs-snsvd- 179 120 TYDNGFIEIHTDKPVYTPDQSVKVRVYSINDDIKPAKRETVITFIDPEGSEVDMVEEIDH 179 180 lnff-wp-ynlpdlvs1gtwrivaky-ehspenytayfdvrkyvlpsfevrlqpsekff- 235 ::: | :::| |:|| | | | | | | | : | ||||| IGIISEPDEKIPSNPRYGMTIKAKYKEDESTTGTAYEVKEYVLPHFSVSIEDEYNFIG 239 62 prkqktlf trvdmnpaggmlvtptieipakevstdsrqnqyvvvqvtgpqvrlekvvll 121 236 yidgnenfhvsitarylygeev-egvafvlfgvk--i-ddakksipdsltripiidgdgk 291 4 malylvaalligfpg-s-shgalytlitpavlrtdteeqilveahgdstpkqldifvhdf 61 and CVF2 may be used in the treatment of cancer by linking them to Cobra; C3; third component of complement; human; mouse; rat; X. laevis; pre-pro molecule; beta chain; alpha chain; codon usage; DNA encoding cobra C3, CVF 1 and CVF 2 - which are used in the G+C content; immune response; host defence; tumour; ss. Bredehorst R, Fritzinger DC, Vogel C; WPI; 94-333186/41. N-PSDB; Q77790. English. selective destruction of cancer cells. Location/Qualifiers R63223 standard; Protein; 1642 AA. R63223; Claim 3; Fig 2W-2J(10); 155pp; Protein 733..1032 /note= "Gamma chain" Protein 1264..1642 /note= "Signal peptide"
Protein 23..1642
Protein 23..649
/note= "Alpha chain" 502; Conservative 30-JUN-1995 (first entry) 07-APR-1993, US-043747. (GEOU) UNIV GEORGETOWN. 07-APR-1994; U03441. treatment of cancer Protein /note= "Beta chain" 1642 AA; WO9423024-A. 13-0CT-1994. Cobra CVF1 Naja naja. Peptide 180 DDT TO THE FETT TO g g 셤 원 셤 ð à ð δ

유

\$

ð

윤

Š

g

Š

셤

õ

g

δ

| : : :: : : : : : : : | Qy 1171 | |
|---|---------------------------------------|---|
| <pre>292 atlkrdt-frs-rfpnlnelvghtlyasvtvmtesgsdmvvteqsgihivaspyqihftk 349 : : :: :: : : : : </pre> | Db Qy | 1203ey-nahthniegtsya : :: :! :: ! : 1231 NLQHKDSSVPNTGTARMVETTAYA |
| 350 tpkyfkpgmpyeltvyvtnp-dgs-paa-hvp-v-vr-eafh-smgtt-ledgta 396 | Db Qy | 1254 vmafqalaeyeiqmpthkdlnldi |
| 397 klilniplnaqslpitvrtnhgdlprerqatksmtaiayqtqggsgnylhvaitsteikp 456 :: : | Db Qy | 1314 tas-gdgkatmtiltfynaqlqek ::: : : 1349 STGFGSGLATWHVTTVVH-KTSTS |
| 457 gdnlpvnfnvkgnanslkqikyftylilnkgkifkvgrqprrdgqnlvtmnlhitpdlip 516 :: :: :: :: | Db Qy | 1370 ry-lgevds-tm-tiidismlt : : ::: : 3XKPSREESSSGSSHAVMDISLPT |
| 517 sfrfvayyqv-gnneivadsvævdvkdtcmgtl-v-vkgdnliqmpgaamkiklegdp 571 : : : :::: : : | Db Qy | 1425 iylnkvshsedeclhfkilkhfev : :: : : : 1 1462 LQLNSIPSSDFLCVRFRIFELFEV |
| 572 garvglvavdkavyvlndkykisqakiwdtieksdfgctagsgqmnlgvfedaglaltts 631 | Db Qy | 1485 cignvcrcagetcsslnhq-e-ri : : : 1520 CEGAACKCVEADCGQMQEELDLTI |
| 632 tnintkqrsaakcpqpanrrrrssvllldsnaskaaefqdqdirkccedvmhenpmgytc 691 | Db Qy | 1543 dvlevikggtdknprakthgyisg :1:: :: i 1580 TLLDIYKTGEAVAEKDSEITFIKK |
| 692 ekrakyiqegdackaafleccryikgvrdenqreselflarddnedgfiadsdiisrsdf 751 | Db Qy | 1601 it-kn-twierwphedecqe-eef : |
| 752 pkswiwitkditeepnsggissktmsfylrdsittwv:lavsftptkgicvaepyeirvm 811 | RES | RESULT 7 ID R63224 standard; Protein; 133 |
| 812 kvffidlgmpysvvkneqveirailhnyvnedi-y-vrvellynpafcsastkg 863 | AC DE DE KW | R63224; 05-JUL-1995 (first entry) Cobra partial CVF2. Cobra; C3; third component of |
| 864 gryrqqfpikalssravpfvivplegglhdveikasvqealw-sdgv-rklkvvpegvq 921 : : :: : :::: : : 881 SKCVRQ-KVEGSSSHLVTFTVLPLEIGLHNINFSL-ET-WFGKEILVYTLRVVPEGVK 935 | KW KW OS | is; pre-p response; ija. |
| 922 ksivtivkldprak-gvggtqlevikarklddrvpdteietkiiiggdpvaqiiensidg 980 : : : : : : : : : | # # # # # # # # # # # # # # # # # # # | "Partial "Gamma ch |
| 981 skinhliitpsgcgeqnmirmaapviaty-yldtteqwetlginrrte-a-vnqivt-gy 1036 | FF P G C | Region 9471333 /note= "Beta chain" W09423024-A. 13-ocT-1994. |
| 1037 aqqmvykkadhsyaaftnrassswltayvvkvfamaakmvaqisheiicqqvrwlilnrq 1096 :: : LSIMSYRNADYSYSVWKGGSASTWLTAFALRVLGQVNKYVBQ-NQNSICNSLMLVENYQ 1112 | PR PA PI | U7-ARK-1994; U03441. 07-ARR-1993; US-043747. GEOU) UNIV GEORGETOWN. Bredehorst R, Fritzinger DC, EDI. 01-33316.//1 |
| 1097 qpdgafkenapvlsgtmqggiqgaeeevyltafilvalleskticndyvnsldssikk 1154 : : : : ::: : ::: : 1113 LDNGSFKENSQYQPIKLQGTLEVEARENSLYLTAFTVIGIRKAFDICP-LVK-IDTALIK 1170 | DR PT | N-PSDB; Q77791. NA-PscBr; Q77791. UNA encoding cobra C3, CVF 1 treatment of cancer C1:5 5: 2: 2: 2: 2: 5: 2: 2: 5: 7: 5: 7: 5: 7: 7: 7: 7: 7: 7: 7: 7: 7: 7: 7: 7: 7: |
| 1155 atnyllkkyeklqrpyttaltayalaaadqlndd-rvlm-aa-stgrdh-w-e 1202 | 3 2 | Claim 3; kig 2M-2V; 133pp; En This sequence represents the |

g

Š

요

8

음

Š

셤

Š

셤

රු පු රු

Š

g

g

ð

g

ð

Š

ekanvcnkfhlnvsvenihlna-mg-akgalmlki-ct 1369 :||:||:|:::| ::: :: :: |: TSEEVCS-FYLKIDTQDIEASHYRGYGNSDYKRIVACA 1406 sqrkcqealnlkvnddyliwgsredllptk-d-kisyi 1600 | i | i | i | i | | i : | | i : | | KKVTCTNA-ELVKGRQYLIMG-K-EALQIKYNFSFRYI 1636 ltgflpdaedltrlskgvdryisryevdnnmaqkvavi 1424 ||:::||| ||||:::|: || PTGISANEEDLKALVEGVDQLFTDYQI-KD-GH---VI 1461 ridvplqiekacetnvdyvyktkllrieeqdgndiyvm 1542 | :: | | :: | | : | | TISAETRKQTACKPENAYKYSITSITVENVFVKYKA 1579 English. e C-terminal portion of cobra venom factor f complement; human; mouse; rat; beta chain; alpha chain; codon usage; 1 and CVF 2 - which are used in the eef-qklcddfaq 1630 :| :| !|!! ?AFLANL-DEFAE 1669 c, Vogel C; ce; tumour. alifiers 333 AA.

888888888888

අ 3 유 à 유 à g Š a Š 용 ð 유 ð 셤 à g ò a 8 셤 õ 유 8

9

8

Z

: | :: : | :: | | 1386 | 1894 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 | 1896 1095 ONQNSICNSLIMLVENYQLDNGSFKENSQYQPIKLQGTLPVEARENSLYLTAFTVIGIRK 1154 1273 WLSEEQRYGGGFYSTQDTINAIEGLTEYSL-LVKQLRLSMDIDVSYKHKGALHNYKMT-D 1330 979 nallaqtvetkinedftvsas-gdgkatmtiltvynaqiredanvcnkfhidvsvenvql 1037 1038 nlkeakgakgalklkictry-lg--evds-tm-tiidvsmltgfvpdtedltrlskgvdr 1092 1389 SHYRGYGNSDYKRIVACASYKPSREESSSGSSHAVMDISLPTGISANEEDLKALVEGVDQ 1448 1093 yismfeinnnmaqkgtviiyldkvshsedeclhfkilkhfevgfiqpgsvkvysyynlde 1152 1504 QCTMFYSTSNIK--IQKVCEGAACKCVEADCGQMQEELDLTISAETRKQTACKPEIAYAY 1561 1211 ktkllrieekdgndiyvmdvlevlkggtdqnqqvkvrqyvsqrkcqealnlmvnndyliw 1270 918 978 705 inrrte-a-vngimt-gyaqqlvykkadhsyaaftnsassswltayvvkifalaakivkd inheivcggmrwlilnrqrtdgvfrenapvlfgtmqggiqgaepegsl--tafilvalle srsicn a yinilds siskat dyllk yeklqrpyttaltayalaaa erlndd-r--v--lm-aa-stgrn---r-w-e---e---pnaht-hniegtsyallallkmkkfveagpvvg 919 wlidqqyyggtygqtqatvmmfqalaeyeiqmpthkdlnlditielpdrevpiryrinye || ::| ||| : || || : ::|| || : : || :: | 1271 gpss-dl-wpmkdkisylitkntwierwphedkcqe-eef-gklcddfa 1315 | : :: | :| |||| ||:: | :| |:||| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||:|| ||: = :: :: :: :: : : - -<u>::</u> <u>-</u> _ :: :: 셤

용

ď

ò

g

õ

셤

ð

рp

à

g

ð 셤 Š В 8

Š

g

g

8

g

S

Š

g

ð

 Ξ

US-08-487-283A-2,rag

1137 1167 1313 : |: : | :: || :: | :: || 1051 || 1051 || 1051 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 || 1055 pqkpkapvghfyepqapsae-vemtsyvllayltaqpaptsedltsatnivkwitkqqna 1254 slpelpgeysmkvt-gegcvylqtslkynilpekeef-pfalgvqtlp-qtcdepk-aht 1369 969 m-qntqnll-qmpygcgeqnmvlfapniyvldylnetqqltpeikska-ig-y-lntgy- 1022 qrqlnykhy-dgsystfgerygrnggntwltafvlktfaqarayifideahitqaliwls 1081 EVL-INDDLIVSTGFGSGLATVHVTTVVHKTSTSEEVCSFYLKIDTQDIEASHYRGYGNS 1397 1370 sf-qislsvsytger--sas-nm-aivdvkmvsgfiplkptvkml-ersnhv-srtevss 1422 ffveltmpysvirgeaftlkatvlnylpkcirvsvqleaspaf-la-vpvekegaphci- 858 882 918 896 706 NNDETCEORAARISLGPRCIKAFTECCVVASQLRANISHKDMQLGRLHMKTLLPVSKPEI 765 801 rkyfpetwiwdlvvvnsagvaevgvtvpdtitewkagafclsedaglgisstaslrafqp RSYFPESWLWEVHLVPRR--KQLQFALPDSLTTWEIQGIGIS-NTGICVADTVKAKVFKD cangrqtvswavtpkslgnvnftvsaealesqelcgtevpsvpehgrkdtvikpllvepe CV--RQKVE-GSSSH-L--VTFTV-L-PLEI-GLHNINPSLETWFG-KEILVKTLRVVPE glekettfns1-lcpsg--gevs--eelslklppnvveesa--ra-sv-svl-gdilgsa qr-qkdngcfrssgsllnnaikggvedevtlsa-yit-ialleipltv-thpvvrnalfc lesawktaqegdhgsh-vytkallayafalagnqdkr-kevlkslneeavkkdnsvhwer LIKADNFLLENTLPAQSTFTLAISAYALSLGDKTHPQFRSIVSALKREALVKGNPPIY-R qggfsstqhtvvalhalskyga-atftrtgkaaqvtiqssgtfsskfqvdnnnrlllqqv 1423 nhvliyldkv-snqtlslfftvlqdvpvrdlkpaivkvydyyetd 1466 Recombinant human alpha-2 macroglobulin <u>::</u> Ė = :: R11334 standard; Protein; 1 R11334; 04-JUN-1991 (first entry)

Jan 28 12:20

US-08-487-283 A-2 rag

| S × | Homo sapiens. AO9103557-A. | đ | 599 avdqsvllmkpdaelsassvynllpel |
|----------------|--|----------|--|
| <u>5</u> بر | 21-WAR-1991. 38-aric-1990 - R6025 | δ | 604 AVDSAVYGVQRGAKKPLERVFQFLEKS |
| t & # | 29-AUC-1930, DNO.222. 29-AUC-1989, DK-004237. 20-AUC-1980 - DK-004237. | qq | 655 -yt-pvsst-nek-dmysfledmglka |
| £ 65. 5 | 29-AUC-1989; DK-04235. 29-AUC-1989; DK-04236. | δ | 664 ENDEPCKEILRPRRTLØKKIEEIAAK- |
| 5 2 2 | (INVO.) NOVO NOMIZON A/J. Esper Lare S-J; Ent. 01 100/15/14 | q | 704 r-v-gfyesdvmgrgh-a-r] |
| 5 55 5 | N-PSDB, Q11581. NN-STBB, Q11581. | δ | 723 RCIKAFTECCVVASQIRANISHKDMQI |
| . 5. 5 | The encoding aight macks ground - used to produce recomming to use in growth media, proteinase inhibitors and as | qq | 749 agvaevgvtvpdtitewkagafclsec |
| - x x | catters Disclosure; page 34; 78pp; English. Phis roomhisant human alpha-2 macroalphulin is magful as an add | Qy | rweig |
| 3 23 5 | into recombinant number appre-z macrogrobusin is useful as an aug- litive to growth-handia, as a proteinase inhibitor and as a carrier | qu | 809 tlkatvlnylpkcirvsvqleaspaf |
| 3 8 | in gene- and enzyme repiacement-therapy. See also vilsol. Sequence 1474 AA; | δ | 840 QLKGTVYNYRTSGMQFCVKMSAVEGIC |
| Önö | Query Match 4.7%; Score 564; DB 2; Length 1474; | QQ | 866 gnvnftvsaealesgelcgtevpsvpe |
| X | t bocal Similarity 22:33, Fred. No. 3:08=50, ches 350; Conservative 408; Mismatches 629; Indels 141; Gaps 118; | δλ | 4 |
| e & | 27 gkpq-ymvlvpsllhtettekgcvll-sylne-tvtvsa-slesvr-gnrslftdleaen 81 :: ::: :: : :: : | do yo | 925gevseelslkippnvveesar : :: :: : 949 IYGTISRRKEFPYRIPLDLVPKTEIKE |
| ą | 82 dvlhcvafav-pks-ssnee-vmfltvqvkgptqefkkrttvmvknedslvfvqtdksiy 138 | QQ | 974 eqnmvlfapniyvldylnetqqltpei |
| ⋧ | : ::: :: :: :: :: : : : : : :: :: | δy | ::: : : :: :: : : : : : : |
| ą | 139 kpgqtvkfryvsmdenfhplneliplvyiqdpkgnriaqwqsfqlegglkqfs-fplsse 197 | QQ | 1029 gerygrnggntwltafvlktfagaray |
| ⋧ | | ٥y | 1067 SVWKG-GSASTWLTAFALRVLGQVNKY |
| ą | 198 pfggsykvvvq-kksggrtehp-ftveefvlpkfevqvtvp-kiitile-eemnvsvcgl 253 | ପ୍ଧ | 1088 nnaikggvedevtl8a-yit-iallei : : : : : : |
| à | 194 PRYCMMTIKAKYKEDESTTGTAYFEVKEYVLPHFSVSIEPEYNFIGYKNFKNFEITIKAR 253 | δy | 1126 PIKLOGTLPVEARENSLYLTAFTVIGI |
| ન ≿ | 254 ytygkpyp-ghvtvsicrkysdasdchgedsq-afcekfsgqlnshgcfyqqvktkvfgl 311 | Ub Qy | 1144 vytkallayafalagnqdkr-kevlke |
| ą | 312 -krkeyemk-lhteaq-iqeegtvveltgrqsseitrtiitklafvkvdshfrqqipffgq 368 | qq | 1203 sae-vemtsyvllayltagpaptsedl |
| ≿ | 313 YSLEDLWNKYLYIAVTVIESTGGFSEEAEIPGIKYVLSPYKLALVATPLELKPGIPYPIK 372 | δλ | : |
| ą | 369 vrlvdgkgvpipnkvifirgneanyysnatt-d-ehglvqfsinttn-vmgtsltvrv-n 424 | QQ | 1262 skyga-atftrtgkaaqvtiqssgtfs |
| ⋩ | LDQLVGGVPVIIMAQTIDVNQ | 0y | IDVSYKHK |
| ą | 425 ykdrspcygywwseeheeahhtaylvfspsksfwhlepmshelpcghtqtvqahyilng 484 | Dp | 1320 gcvylqtslkynilpekeef-pfalgv |
| à | 433 FNVKTDAPDLPEENQAREGYRALAYSSLSQSYLYIDWTDNHKALLVGE-H-LNI-IVTPK 489 | δ | 1355 GLATVHVTTVVHKTSTSEEVCSFYLKI |
| ą | 485 gtllglkklsfyylimakggivrtgthgllvkqedmkghfsisipvksdiapvarlliy- 543 : : : :: : : : : : : | QQ | 1374 sas-nm-aivdvkmvsgfiplkptvkm |
| ⋧ | 490 SPYID-KITHYNYLILSKGKIIHFGTREKFSDASYQ-SINIPVTQNMVPSSRLLVYY 544 | ογ | 1415 SSSGSSHAVMDISLPTGISANEEDLKA |
| ያ ጽ | 544 avlptgdvigdsakydven-clankvdlsfsp-sqslpasha-hlrvtaapqsvcalr 598 ::::: :: : :: : ::: 545 IVTGEQTAELVSDSVWINIEEKC-GNQLQVHLSPDADAYSPQGTVSLNMATGMDSWVALA 603 | d y | 1429 lfftvlqdvpvrdlkpaivkvydyyet |
| | | | |

Š

셤

g

õ

õ 셤 Ş 용 Š g Š 셤 Š 셤 Š

음

g

Š

9

õ

eikska-ig-y-lntgy-grqlnykhy-dgsystf 1028 :|:|:||:||: IFHSDPLIEKQKLKKKLKEGMLSIMSYRNADYS-Y 1066 kml-ersnhv-srtevssnhvliyldkv-snqtls 1428 | | | ::: : : | |:: | :: | KALVECVDQLFTDYQIKDGHVILQLMSIPSSDFLC 1474 ayifideahitqaliwlsqr-qkdngcffssgsll 1087 |: :: | :|:||: | ||| :: KYVEQNQNSICNSLLWLVENYQLDNGSFKENSQYQ 1125 iltsatnivkwitkqqnaqqqfsstqhtvvalhal 1261
!: ::||:::| || || || ||::|
>INYVNPVIKWLSEEQRYGGGFYSTQDTINAIEGL 1297 sskfqvdnnnrlllqqvslpelpgeysmkvt-ge 1319 yqtlp-qtcdepk-ahtsf-qislsvsytgsr-- 1373 ::| :: ::: :| || || KIDTQDIEASHYRGYGNSDYKRIVACASYKPSREE 1414 silltv-thpvvrnalfclesawktaqegdhgsh- 1143 kslneeavkkdnsvhwerpqkpkapvghfyepqap 1202 :|: ||: | |: | SALKREALVKGNPPIY-RFWKDNLQHKDSSVPNTG 1243 edaglgisstaslrafqpffveltmpysvirgeaf 808 ::|: :: | :: | |:||:|||: -NTGICVADTVKAKVFKDVFLEMNIPYSVVRGEQI 839 703 722 x1--vhvee--phtet-vrkyfaetwiwdlvvvns 748
:| :|: | : :| ||:|::: :|
QLGRLHMKTLLPVSKPEIRSYFPESWLWEVHLVPR 782 ekdlt-gfpgplndqdd-e-dcin-rhnvyingit 654 f-la-vpvekeqaphci-cangrqtvswavtpksl 865 : : || |: | || |: | pehgrkdtvikpllvepeglekettfnsl-lcpsg 924 kaftnskirkp--k-mcpqlqq-yemhgpe-gl--| ::| ::| :: | :: | K-YKHSVVKKCYDGACVNNDETCEQRAARISLGP KSDLGCGAGGGLNNANVFHLAGLTFLTNANADDSQ td 1456 PD 1502

52

Query Match 4.5%; Best Local Similarity 100.0%; 12-SEP-1994; U09770. bond 253 403 AA; WPI; 95-123385/16. Modified site Bredehorst R, WO9507299-A. 16-MAR-1995. Naja naja. Disulfide Sequence R71933; chain" Matches Matches 셤 g ð ð 셤 ð 셤 쇰 õ õ 셤 ð ö 678 TLOKKIEEIAAKYKHSVVKKCCYDGACVNNDETCEQRAARISLGPRCIKAFTECCVVASQ 737 1 tlqkkieeiaakykhsvvkkccydgacvnndetceqraarislgprcikafteccvvasq 60 0; Gaps C5a receptor antagonists having no agonist activity - are used in compsns. to treat C5a-mediated diseases and inflammatory conditions Disclosure; Page 36-37; 65pp; English. C5a encoded by a synthetic gene is given in R75497. Analogs of C5a, obtd. by mutagenesis of the C-terminal region, exhibit excellent antagonistic properties and substantially no agonist activity. The synthetic gene encoding this protein was constructed from ten 41-mer oligonuleotides. The sequence was chosen to maximise E.coli E.coli is posssible by inclusion of an ATG codon in front of the triplet coding for the first amino acid of the C5a polypeptide. See also N70213-6, N70218, N71339, N71340, N71343-8. Davidow LS, Franke AE, Dezeeuw JR. WPI; 87-124409/18. New Yarrowia lipolytica transformants - used fro expression and secretion of heterologous proteins, esp. prorennin, and human and S.cerevisiae preferred codon utilisation and to allow for several restriction enzyme sites. Direct expression of C5a in Score 541; DB 2; Length 74; Pred. No. 5.04e-34; 0; Indels Peppard JV, Van Oostrum J; Human C5a protein. C5a; C5a receptor-antagonist; antiinflammatory. 0; Mismatches activated complement component C5a P71666 standard; Protein; 74 AA. Example; Fig 9; 45pp; English. R75497 standard; Protein; 74 Query Match 4.5%; Best Local Similarity 100.0%; 25-APR-1991 (first entry) 74; Conservative 13-NOV-1995 (first entry) Boyar WC, Galakatos NG, WPI; 95-224319/29. Human anaphylatoxin C5a. 10-OCT-1986; 307839. 18-OCT-1985; US-789206. 18-MAR-1986; US-841121. 06-DEC-1993; US-162591. 61 lranishkdmqlgr 74 (CIBA) CIBA GEIGY AG. 16-NOV-1994; IB0359. (PFIZ) PFIZER INC. anaphylatoxin C5a 74 AA; N-PSDB; Q92518. EP-220864-A. W09516033-A. 06-MAY-1987. 15-JUN-1995 Synthetic. Synthetic. Sequence Sequence P71666; R75497; Matches RESULT

셤 ð 용 à ö 175 dnnmaqkvavijylnkvshsedeclhfkilkhfevgfiqpgavkvysyynlderctkfyh 234 :: :: || :: || :| || :| || 11156 -KD-GH---VILQIANSIPSSDFLCVRFRIFELFEVGFLSPATFTVYEYHRPDKQCTWFYS 1510 64 vetklnqditvtas-gdgkatmtiltfynaqlqeranvcnrphlnvsvenihlna-hg-a 120 121 kgalmlki-ctry-lg--evds--thtiidismltgflpdaedltrlsrgvdryisryev 174 1 tlqkkieeiaakykhsvvkkccydgacvnndetceqraarislgprcikafteccvvasq 60 4 ygetyggtgatvmafgalaeyeigmpthkdlnlditielpdrevpiryrinyenallart 63 Gaps 112; Conservative 105; Mismatches 156; Indels 32; Gaps used for activating complement partic. in killing tumour cells Disclosure; Fig. 9; 36pp; English. New cobra venom factor (CVF) derivs. are obtained by chymotrypsin or trypsin digestion of the C-terminal region of CVF beta-chain Grunwald T, Ollert MW, Vogel C, Ziegelmuller P; (given in R71933). The CVF derivs. retain complement activating New cobra venom factor deriv. - produced by protease digestion, ; activity but are less prone to neutralization by the immune Score 541; DB 13; Length 74; Pred. No. 5.04e-34; Query Match 4.5%; Score 540; DB 13; Length 403; Best Local Similarity 27.7%; Pred. No. 6.15e-34; 0; Indels /note= "interchain disulfide linkage to CVF gamma 0; Mismatches Cobra venom factor C-terminal region. Cobra venom factor; CVF; cytostatic; tumor. Location/Qualifiers .T 12 R71933 standard; protein; 403 AA. /label= N-glycosylation_site 21-SEP-1995 (first entry) Conservative response than natural CVF. 738 LRANISHKDMQLGR 751 (GEOU) UNIV GEORGETOWN 61 lranishkdmqlgr 74 10-SEP-1993; US-118674.

```
98; Mismatches 127; Indels 28; Gaps 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      69 ldqteqwekf-g--lekrqealelikkgytqqlafkqpssayaafnnrppstwltayvvk 125 | : ::| | : | :| | :| | :| | 1084 | 1084 | 1085 lETGNHWNIFHSDPLIEKQKIKKKIKEGMLSIMSYRNADYSYSVWKGGSASTWLTAFALR 1084
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   126 vfslaanliaidsqvlcgavkwlilekqkpdgvfqedgpvihqemiggfr-ntkeadvsl 184
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           185 tafvlialqeardicegqvnslpgsinkagetleasylnlqrpytvaiagyalalmnkle 244
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Prodn. of 14kDaII type phospholipase A2 inhibitor - by treating specified proteins with organic and/or acidic solvent, to obtain enzyme useful to treat inflammation and allergy claim 1; Page 5-6; Bpp; Japanese.

A 14kDaII type phospholipase A2 inhibitor, (G3dg) is produced by treating the polypeptide R31948 with an organic and/or acid solvent. Rat C3dg can be mass-produced by genetic engineering for use in the treatment of inflammation and allergic diseases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10 dqvpdtdsetril-lqgtpvaqmaedavdgerlkhlivtpsgcgeqnmigmtptviavhy 68
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match 3.6%; Score 429; DB 10; Length 344; Best Local Similarity 28.5%; Pred. No. 1.78e-24; Matches 101; Conservative 98; Mismatches 127; Indels 28
                                                                                                                                                                Phospholipase A2 inhibitor - C3dg (Rat).
Phospholipase A2; Inhibitor; Rat C3dg; production;
organic solvent; acid solvent; Allergy; Inflammation.
.r 15
R51948 standard; protein; 344 AA.
                                                                                                                          08-NOV-1994 (first entry)
                                                                                                                                                                                                                                                                                       Ratus sp.
J0665293-A.
08-MAR-1994.
21-AUG-1992; J22693.
11-AUG-1992; JP-222693.
(TEIJ ) TEIJIN LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WPI; 94-121221/15.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Sequence 344 AA;
                                                                                        R51948;
   RESULT ID RESULT                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              പ്പ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ð
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             g
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              à
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      q
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Š
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           셤
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        q
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Š
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ð
```

Search completed: Wed Jan 28 12:13:50 1998 Job time : 66 secs.

ð 셤 õ g ð Rattus rattus WO9202619-A. 20-FEB-1992. 17;

18

Suzuki Y;

89 Gaps

-::

See also R21776.

Sequence

셤

셤

Š 셤 õ

ð

184

= = = = =

= - :::: =

185

셤